NAVAL POSTGRADUATE SCHOOL Monterey, California



THESIS

DEVELOPMENT AND IMPLEMENTATION OF A SHELL ELEMENT WITH PRESSURE VARIATION THROUGH THE THICKNESS AND VOID GROWTH AND NUCLEATION EFFECTS

by Patrick M. McDermott

September 1999

Thesis Advisor:

Young W. Kwon

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REPORT DOCUMENTATION PAGE

Form Approved OMB No. 0704-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instruction, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Washington headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reduction Project (0704-0188) Washington DC 20503.

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1. AGENCY USE ONLY (Leave blank)	2. REPORT DATE September 1999		TYPE AND DATES COVERED al Engineer's Thesis	
4. TITLE AND SUBTITLE DEVELOPMENT AND IMPLEMENTATION OF A SHELL ELEMENT WITH PRESSURE VARIATION THROUGH THE THICKNESS AND VOID GROWTH AND NUCLEATION EFFECTS			5. FUNDING NUMBERS	
6. AUTHOR(S) McDermott, Patrick M.				
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) Naval Postgraduate School Monterey, CA 93943-5000		8. PERFORMING ORGANIZATION REPORT NUMBER		
9. SPONSORING / MONITORING AGENCY NAME(S) AND ADDRESS(ES)		10. SPONSORING / MONITORING AGENCY REPORT NUMBER		
11. SUPPLEMENTARY NOTES				
The views expressed in this thesis of the Department of Defense or	the U.S. Government.	not reflect	the official policy or position	
12a. DISTRIBUTION / AVAILABILITY STATEMENT		12b. DISTRIBUTION CODE		
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13. ABSTRACT (maximum 200 words)

A shell formulation was developed from a three-dimensional solid. The shell element has four corner nodes at which there are three displacements and three rotations as nodal degrees of freedom, and includes both transverse shear and transverse normal deformations. The element utilizes reduced integration along the in-plane axes and full integration along the transverse axis. The formulation incorporates the Gurson constitutive model for void growth and plastic deformation. An algorithm for stable solutions of the nonlinear constitutive equations is also developed. Hourglass mode control is provided by adding a small fraction of internal force determined through full integration along the in-plane axes and reduced integration along the transverse axis. Implementation into a research finite element program is discussed. Numerical examples are provided to verify the accuracy of the new element and to show the importance of the transverse normal stress, void effects on plastic strain, and the necessity of applying a drilling moment.

14. SUBJECT TERMS Finite Element, Shell Formulation, Void Model, Elasto-plastic, High Order Element			15. NUMBER OF PAGES
17. SECURITY 18. SECURITY CLASSIFICATION OF 10. DECURITY OF ACCURATION.			16. PRICE CODE
17. SECURITY CLASSIFICATION OF REPORT Unclassified	THIS PAGE Unclassified	19. SECURITY CLASSIFI- CATION OF ABSTRACT Unclassified	ABSTRACT UL

NSN 7540-01-280-5500 239-18

Standard Form 298 (Rev. 2-89)

Prescribed by ANSI Std.

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DEVELOPMENT AND IMPLEMENTATION OF A SHELL ELEMENT WITH PRESSURE VARIATION THROUGH THE THICKNESS AND VOID GROWTH AND NUCLEATION EFFECTS

Patrick M. McDermott Lieutenant, United States Navy B.S.E.E., Memphis State University, 1993

Submitted in partial fulfillment of the requirements for the degree of

MECHANICAL ENGINEER

from the

NAVAL POSTGRADUATE SCHOOL September 1999

Author:

Patrick M. McDermott

oung W. Kwon, Thesis Advisor

Approved by:

M. D. Kelleher, Acting Chair

Department of Mechanical Engineering

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I. INTRODUCTION

Since shell structures are efficient load-carrying structural members, they have been dominant in most structural applications. However, the finite element formulation of shells poses some difficulty. As a result, extensive study has been devoted to developing better shell elements. Because of the abundance of papers on this subject, no attempt is made here to summarize all of them. Some of the related past work is given in references [1-12].

Void growth and nucleation can have a significant effect on plastic flow [13]. Since voids act as stress concentrators, the overall effect is to reduce the stress under plastic flow, and increase the plastic strain [14-16]. The model proposed by Gurson [17] appears to have been adopted as the standard for incorporating void growth and nucleation effects into a numerical solution of elasto-plastic problems. Previous work has been done on improving the efficiency and accuracy of plasticity computations, applying plasticity to plate/shell elements, and incorporating void growth and nucleation effects in solid elements [18-20]. The element presented in this paper incorporates a modified version of the algorithm for three-dimensional solids proposed by Aravas [21] into a shell element. This shell element assumes a modified plane-stress condition, and utilizes both the hydrostatic pressure and the deviatric stress. Due to the importance of the hydrostatic pressure on void growth and nucleation, this element includes the transverse normal stress.

The algorithm proposed by Aravas is not unconditionally stable when applied to this modified plane stress condition [21]. This paper also presents modifications to the original algorithm, which greatly enhance the stability of the solution, at the cost of a slight decrease in computational efficiency. This algorithm also allows for more complicated work-

hardening profiles than the typical exponent law ($\sigma = K\epsilon^n$). Full modeling of the entire stress-strain curve is accomplished by a piece-wise linear approximation. While this method is not particularly useful for analytic approaches, it appears to be helpful in strictly numerical solutions. The stress-strain relationship may be taken directly off the results of a standard tensile test.

For thick shell applications, the transverse normal stress and strain cannot be neglected. The transverse normal stress and strain affect both the hydrostatic pressure and the deviatoric stress, which in turn affect plastic deformation and void growth and nucleation. The element presented here extends the typical use of the drilling degree of freedom (DOF), as outlined in Hughes and Brezzi [22], to incorporate the effects of transverse normal strain. The drilling DOF is important in transmitting stress and strain to elements across sharp bends and curves, and is therefore already included in a variety of shell elements [2,4,6,7,10]. In addition to this traditional usage, the present element utilizes the drilling DOF to compute the transverse normal deformation. The importance of including the transverse normal deformation is outlined in Essenburg [23].

The present study formulates a shell element for transient analysis. The element can have elasto-plastic deformation with void growth and nucleation. Gurson's void model is used as a basis for the void constitutive model. The shell element includes both transverse shear deformation and the transverse normal deformation for thick shell applications. The drilling degree of freedom is used for computing the deformation through the thickness of

a thick shell. An algorithm for stable solutions of the nonlinear constitutive equations is also developed.

Some example problems are presented to evaluate the formulation and to investigate the effects of the transverse normal strain for thick shells in association with elasto-plastic deformation, including void effects. The sections that follow are: the formulation of the shell element, discussion of the elasto-plastic constitutive equation including void nucleation and growth, the stable solution algorithm including spurious (hourglass) mode control, numerical examples and discussion of results, and conclusions.

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II. FINITE ELEMENT FORMULATION

A. GEOMETRY

A point in a shell structure can be expressed by a vector sum of two vectors. The first vector is a position vector from the origin of the global coordinate system to a point on a reference surface of the shell element. The second vector is a position vector from this reference surface to the point under consideration. The surface that spans the center of the transverse axis is used as the reference surface in this formulation, although any surface would suffice. The first vector terminates at the reference surface directly below the point in question. The second vector is then the normal from the reference surface that intersects the desired point. Figure 1 shows this relationship. Two shape functions are used to describe

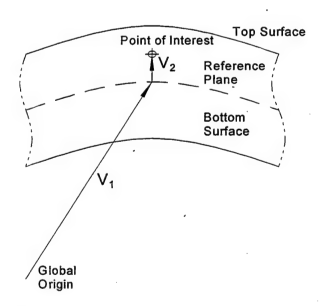


Figure 1. Element Cross Section.

a position in the element; N^k is the two dimensional shape function in the ξ - η plane, and H^k is the one dimensional shape function along the ζ axis, where (ξ, η, ζ) describes a point in the natural coordinate system. A generic point in the shell may now be described in terms of the position vectors of the nodes and the shape functions:

$$x_{i}(\xi,\eta,\zeta) = \sum_{k=1}^{n} N^{k}(\xi,\eta) \ x_{i}^{k} + \sum_{k=1}^{n} N^{k}(\xi,\eta) H^{k}(\zeta) V_{3i}^{k} \ (i = 1,2,3)$$
 (1)

where x_i^k is the position vector of node k in the reference surface; V_{3i}^k is the unit vector at the node k; and n is the number of nodes per element. In the present formulation, a four-node shell element is considered. The unit vector V_{3i}^k is defined as:

$$V_{3i}^{k} = \frac{(x_{i}^{k})^{top} - (x_{i}^{k})^{bottom}}{\|(x_{i}^{k})^{top} - (x_{i}^{k})^{bottom}\|}$$
(2)

where top and bottom indicate the top and bottom surfaces of the shell, and || || denotes the Euclidean norm. The one-dimensional shape function H^k is expressed as:

$$H^{k}(\zeta) = \left[\frac{1}{4} (1 + \zeta)(1 - \overline{\zeta}) - \frac{1}{4} (1 - \zeta)(1 + \overline{\zeta}) \right] \| (x_{i}^{k})^{top} - (x_{i}^{k})^{bottom} \|$$
 (3)

in which $\overline{\zeta}$ indicates the location of the reference surface and varied from -1 to 1 ($\overline{\zeta} = 0$ denotes the mid-surface). The two-dimensional shape function N^k is expressed as:

$$N^{1} = \frac{1}{4}(1-\xi)(1-\eta)$$

$$N^{2} = \frac{1}{4}(1+\xi)(1-\eta)$$

$$N^{3} = \frac{1}{4}(1+\xi)(1+\eta)$$

$$N^{4} = \frac{1}{4}(1-\xi)(1+\eta)$$
(4)

B. DISPLACEMENT

The displacement field in a shell can be written as:

$$u_{i}(\xi,\eta,\zeta) = \sum_{k=1}^{n} N^{k}(\xi,\eta) u_{i}^{k} + \sum_{k=1}^{n} N^{k}(\xi,\eta) H^{k}(\zeta) (-V_{2i}^{k}\theta_{1}^{k} + V_{1i}^{k}\theta_{2}^{k} + V_{3i}^{k}\theta_{3}^{k}) \quad (i = 1,2,3)$$
(5)

in which u_i is the displacement along the x_i axis, u_i^k is the nodal displacement at the node k, and unit vectors V_{1i}^k and V_{2i}^k lie along the reference surface. V_{1i}^k , V_{2i}^k and V_{3i}^k are mutually perpendicular. θ_1^k , θ_2^k and θ_3^k are rotational degrees of freedom along the unit vectors V_{1i}^k , V_{2i}^k and V_{3i}^k , respectively. The right-hand rule is assumed for the positive direction of each rotation. Figure 2 illustrates the relationship among these vectors.

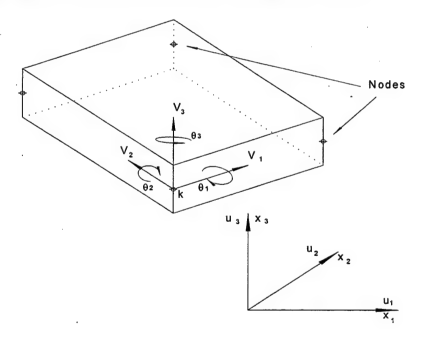


Figure 2. Displacement Vector Orientation.

 θ_1^k and θ_2^k are the bending rotations, while θ_3^k is the drilling rotational degree of freedom. The role of θ_3^k can be clarified by considering a flat plate parallel to the x_1 - x_2 plane. Now Eq. (5) can be rewritten for the transverse displacement as:

$$u_3 = \sum_{k=1}^n N^k u_3^k + \sum_{k=1}^n N^k H^k \theta_3^k$$
 (6)

Equation (6) demonstrates that the transverse deformation varies through the plate thickness (i.e. along the ζ axis) with θ_3^k . In this way, the transverse normal deformation is included in this formulation, along with the transverse shear deformations.

C. COORDINATE TRANSFORMATION

Combining the three unit direction vectors into matrix $[T_p]$ provides the rotation transformation matrix, or matrix of direction cosines, as shown in Eq. (7). $[T_p]^{-1}$ is used to transform the nodal degrees of freedom from the global coordinate system to local coordinates, as shown in Eq. (8), where k is the node number. The components of $\{d\}$ at the four nodes of an element are shown in Eq. (9). Once the internal force vector is generated in local coordinates, $[T_p]$ is used to transform the local vectors back into global coordinates. This procedure will be discussed later.

$$\left[\mathbf{T}_{\mathbf{p}}\right] = \left|\left[\vec{\mathbf{V}}_{1}\vec{\mathbf{V}}_{2}\vec{\mathbf{V}}_{3}\right]_{(3\mathbf{x}3)}\right] \tag{7}$$

$$\{d\} = \{d^1 d^2 d^3 d^4\}^T$$
 (9)

The strain transformation matrix, [T], is used to transform calculated strain from the global coordinate system to local coordinates. Transforming the resulting stress from local coordinates to the global coordinate system would normally require using [T]⁻¹, but since [T] is orthogonal, [T]⁻¹ = [T]^T, where [T]^T is the transpose of [T]. This property negates the requirement to invert a six by six matrix. For a detailed derivation of these transformations, refer to Cook [24]. The strain transformation matrix is explicitly defined in Eq. (10), where V_{ij} is the cosine of the direction vector V_{ij} in the x_{ij} direction.

$$[T] = \begin{bmatrix} V_{11}^2 & V_{12}^2 & V_{13}^2 & V_{11}V_{12} & V_{12}V_{13} & V_{11}V_{13} \\ V_{21}^2 & V_{22}^2 & V_{23}^2 & V_{21}V_{22} & V_{22}V_{23} & V_{21}V_{23} \\ V_{31}^2 & V_{32}^2 & V_{33}^2 & V_{31}V_{32} & V_{31}V_{32} & V_{32}V_{33} & V_{31}V_{33} \\ 2 V_{11}V_{21} & 2 V_{12}V_{22} & 2 V_{13}V_{23} & V_{11}V_{22} + V_{21}V_{12} & V_{12}V_{23} + V_{22}V_{13} & V_{13}V_{21} + V_{23}V_{11} \\ 2 V_{21}V_{31} & 2 V_{22}V_{32} & 2 V_{23}V_{33} & V_{21}V_{32} + V_{31}V_{22} & V_{22}V_{33} + V_{32}V_{23} & V_{23}V_{31} + V_{33}V_{21} \\ 2 V_{11}V_{31} & 2 V_{12}V_{32} & 2 V_{13}V_{33} & V_{11}V_{32} + V_{31}V_{12} & V_{12}V_{33} + V_{32}V_{13} & V_{13}V_{31} + V_{33}V_{11} \end{bmatrix}$$
 (10)

D. STRAIN DISPLACEMENT RELATION

The six components of the strain tensor are computed from Eq. (5) by taking its derivative with respect to the x_i axis. In matrix form, the result for a four-node element is:

$$\{\varepsilon\} = [\mathbf{B}]\{d\} \tag{11}$$

$$\{\varepsilon\} = \{\varepsilon_{11} \varepsilon_{22} \varepsilon_{33} \gamma_{12} \gamma_{23} \gamma_{13}\}^T$$
 (12)

where

$$[\mathbf{B}] = [[\mathbf{B}^1][\mathbf{B}^2][\mathbf{B}^3][\mathbf{B}^4]]$$
 (13)

The detailed expression for $[\mathbf{B}^k]$ is:

$$\left[\mathbf{B}^{k} \right] = \begin{bmatrix} \frac{\partial N^{k}}{\partial x_{I}} & 0 & 0 & -g_{I}^{k} V_{2I}^{k} & g_{I}^{k} V_{II}^{k} & g_{I}^{k} V_{3I}^{k} \\ 0 & \frac{\partial N^{k}}{\partial x_{2}} & 0 & -g_{2}^{k} V_{22}^{k} & g_{2}^{k} V_{I2}^{k} & g_{2}^{k} V_{32}^{k} \\ 0 & 0 & \frac{\partial N^{k}}{\partial x_{3}} & -g_{3}^{k} V_{23}^{k} & g_{3}^{k} V_{I3}^{k} & g_{3}^{k} V_{33}^{k} \\ \frac{\partial N^{k}}{\partial x_{2}} & \frac{\partial N^{k}}{\partial x_{I}} & 0 & -g_{2}^{k} V_{2I}^{k} - g_{I}^{k} V_{22}^{k} & g_{2}^{k} V_{II}^{k} + g_{I}^{k} V_{I2}^{k} & g_{2}^{k} V_{3I}^{k} + g_{I}^{k} V_{32}^{k} \\ 0 & \frac{\partial N^{k}}{\partial x_{3}} & \frac{\partial N^{k}}{\partial x_{2}} & -g_{k}^{k} V_{22}^{k} - g_{k}^{k} V_{23}^{k} & g_{k}^{k} V_{I2}^{k} + g_{k}^{k} V_{I3}^{k} & g_{k}^{k} V_{32}^{k} + g_{k}^{k} V_{33}^{k} \\ \frac{\partial N^{k}}{\partial x_{3}} & 0 & \frac{\partial N^{k}}{\partial x_{I}} & -g_{3}^{k} V_{2I}^{k} - g_{I}^{k} V_{23}^{k} & g_{3}^{k} V_{II}^{k} + g_{I}^{k} V_{I3}^{k} & g_{3}^{k} V_{3I}^{k} + g_{I}^{k} V_{33}^{k} \end{bmatrix}$$

$$(14)$$

in which

$$g_{i}^{k} = \frac{\partial N^{k}}{\partial r_{i}} H^{k} + N^{k} \frac{\partial H^{k}}{\partial r_{i}}$$
(15)

The vector $\{d^k\}$ is defined as:

where u_i is the displacement along the x_i direction at node k, and Θ_i^k is the rotational displacement about the global x_i axis at node k. The matrix $[\mathbf{B}^k]$ must be calculated for each integration point.

E. JACOBIAN MATRIX

Computing the derivatives $\frac{\partial N^k}{\partial x_i}$ and $\frac{\partial H^k}{\partial x_i}$ requires the Jacobian matrix, defined as

$$[\mathbf{J}] = \begin{bmatrix} x_{I,\xi} & x_{2,\xi} & x_{3,\xi} \\ x_{1,\eta} & x_{2,\eta} & x_{3,\eta} \\ x_{1,\zeta} & x_{2,\zeta} & x_{3,\zeta} \end{bmatrix}$$
(17)

where

$$x_{i,\xi} = \frac{\partial x_i}{\partial \xi} = \sum_{k=1}^n \frac{\partial N^k}{\partial \xi} x_i + \sum_{k=1}^n \frac{\partial N^k}{\partial \xi} H^k V_{3i}^k \quad (i = 1, 2, 3)$$
 (18)

$$x_{i,\eta} = \frac{\partial x_i}{\partial \eta} = \sum_{k=1}^n \frac{\partial N^k}{\partial \eta} x_i + \sum_{k=1}^n \frac{\partial N^k}{\partial \eta} H^k V_{3i}^k \quad (i = 1, 2, 3)$$
 (19)

$$x_{i,\zeta} = \frac{\partial x_i}{\partial \zeta} = \sum_{k=1}^n N^k \frac{\partial H^k}{\partial \zeta} V_{3i}^k \quad (i = 1, 2, 3)$$
 (20)

[R] is defined as the inverse of the Jacobian matrix, [J]⁻¹. Then the required partial derivatives are defined as:

$$\frac{\partial N^k}{\partial x_i} = R_{il} \frac{\partial N^k}{\partial \xi} + R_{i2} \frac{\partial N^k}{\partial \eta} \quad (i = 1, 2, 3)$$
 (21)

$$\frac{\partial H^k}{\partial x_i} = R_{i3} \frac{\partial H^k}{\partial \zeta} \quad (i = 1, 2, 3)$$
 (22)

F. STRESS-STRAIN RELATIONSHIP

The strain calculated in Eq. (11) is in the global coordinate, and is transformed to a local coordinate system using

$$\{\varepsilon_{local}\} = [T]\{\varepsilon_{global}\}$$
(23)

where [T] is defined in Eq. (10). Stress is calculated from the strain in the local coordinate system using the plane-strain assumption for the in-plane stress components:

$$\sigma_{x} = \frac{E}{l - v^{2}} (\varepsilon_{x} + v \varepsilon_{y}) \qquad \tau_{xy} = G \gamma_{xy}$$

$$\sigma_{y} = \frac{E}{l - v^{2}} (v \varepsilon_{x} + \varepsilon_{y}) \qquad \tau_{yz} = K G \gamma_{yz}$$

$$\sigma_{z} = E \varepsilon_{z} \qquad \tau_{xz} = K G \gamma_{xz}$$
(24)

where K is the shear correction factor, E is the elastic modulus, G is the shear modulus, and v is Poisson's ratio. The resulting stresses are in the local coordinate system and are converted to the global coordinate system with

$$\left\{\sigma_{global}\right\} = \left[\mathbf{T}\right]^{\mathrm{T}}\left\{\sigma_{local}\right\} \tag{25}$$

$$\{\sigma\} \equiv \{\sigma_{x} \sigma_{y} \sigma_{z} \gamma_{xy} \gamma_{yz} \gamma_{xz}\}$$
 (26)

where [T] is from Eq. (10).

G. DRILLING MOMENTS

Let u_3 be the transverse deflection as defined in Eq. (6). The work done by a pressure loading, p, on an element is:

$$W = \int_{A^c} u_3 \ p \ dA \tag{27}$$

where A^e is the element area. Substituting Eq. (6) into Eq. (27), the work is now:

$$W = \{u_3\}^T \int_{\mathcal{A}} [\mathbf{N}] p \ dA + \{\theta_3\}^T \int_{\mathcal{A}} [\mathbf{N}^k \mathbf{H}^k] p \ dA$$
 (28)

The first term gives the conventional forces at each node, while the second term yields the new nodal load, which will be called the Drilling Moment (DM). For example, if p is a concentrated force P at node n, then the drilling moment associated with θ_3^n becomes $\frac{1}{2}\zeta tP$, where t is the shell thickness and the mid-plane is the reference plane.

When P is applied at the top plane, ζ equals one, and the drilling moment is $\frac{1}{2}tP$. If P is applied at the bottom plane (still with a positive loading direction), the drilling moment is $-\frac{1}{2}tP$. That is, the load results in compression or tension in the transverse normal stress depending on whether the loading is applied to the top or bottom surface of the shell. The transverse normal stress is assumed constant through the shell thickness for elastic deformation. However, as plastic deformation progresses, the transverse normal stress becomes non-uniform through the thickness in order to satisfy the yield function.

The drilling moment direction coincides with the direction of drilling rotation, but affects the transverse normal stress. The drilling moment is used as a mathematical convenience, and thus, lacks a physical interpretation.

H. INTERNAL FORCE, MASS AND THEIR ASSEMBLY

The stress resulting from Eq. (25) is then converted to an internal force vector and summed over all integration points using

$$\{ f_{int} \} = \int_{-l}^{l} \int_{-l}^{l} \int_{-l}^{l} [\mathbf{B}]^{T} \{ \sigma \} d\xi d\eta d\zeta = \sum_{i=1}^{nx} \sum_{j=1}^{ny} \sum_{k=1}^{nz} [\mathbf{B}]^{T} \{ \sigma \}_{\mathbf{W}_{i} \mathbf{W}_{j} \mathbf{W}_{k}} |\mathbf{J}|$$
 (29)

where $|\mathbf{J}|$ is the determinant of the Jacobian matrix, nx, ny, and nz are the number of integration points in the ξ , η , and ζ directions, respectively, and w_i , w_j , and w_k , are the Gauss weights related to those integration points. The resulting force vector, $\{f_{int}\}$, must have its components transformed back to the global coordinate system using:

A lumped mass method is used for the element mass matrix. The matrix for each element is diagonal, with equal diagonal elements:

$$[\mathbf{M}_{e}] = \begin{bmatrix} \mathbf{m}_{e} & 0 & 0 & \dots & 0 \\ 0 & \mathbf{m}_{e} & 0 & \dots & 0 \\ 0 & 0 & \mathbf{m}_{e} & \dots & 0 \\ \dots & \dots & \dots & \dots & \dots \\ 0 & 0 & 0 & \dots & \mathbf{m}_{e} \end{bmatrix}$$
 (31)

where

$$m_e = \frac{\left(\sum_{i=1}^{nx} \sum_{j=1}^{ny} \sum_{k=1}^{nz} w_i w_j w_k |\mathbf{J}|\right) \rho}{n}$$
(32)

with n = 4 in this four-node element. Using a diagonal mass matrix greatly simplifies time integration, since inverting it is a trivial task requiring little computation time. The internal force vector and element mass vector (each 24 by 1) are then assembled into the corresponding system vectors.

I. EXPLICIT TIME INTEGRATION

The use of internal force vectors and explicit time integration negates the need to explicitly form the system stiffness matrices. The acceleration vector is computed from

$$\left\{ \ddot{U} \right\} = \left[\mathbf{M} \right]^{-1} \left(\left\{ F_{ext} \right\}^{t} - \left\{ F_{int} \right\}^{t} \right) \tag{33}$$

where $\{\dot{U}\}$ is the system acceleration vector, [M] is the system mass matrix, $\{F_{ext}\}$ is the system external force vector, and superscript t denotes the time step. Of course, the system mass matrix in Eq. (33) is simply symbolic, since a system mass vector, formed from the diagonal of the mass matrix, is used in actual computation. Velocity and displacement are then found using

$$\left\{ \dot{U} \right\}^{t+\frac{M}{2}} = \left\{ \dot{U} \right\}^{t-\frac{M}{2}} + \left\{ \dot{U} \right\}^{t} \Delta t$$

$$\left\{ U \right\}^{t+\Delta t} = \left\{ U \right\}^{t} + \left\{ \dot{U} \right\}^{t+\frac{M}{2}} \Delta t$$

$$(34)$$

III. DAMAGE CONSTITUTIVE EQUATIONS

A. GURSON'S VOID MODEL

Yielding and plastic deformation in the element follows the model proposed by Gurson for symmetric deformations around a spherical void [17]. The yielding condition is

$$F = \left(\frac{q}{\sigma_0}\right)^2 + 2q_1 \Phi \cosh\left(-\frac{3}{2}\frac{q_2 p}{\sigma_0}\right) - \left(1 + q_3 \Phi^2\right) = 0$$
 (35)

where Φ is the current porosity, p is the hydrostatic stress, q is the effective stress, and σ_0 is the current yield stress. This model assumes equivalent yield stress in both tension and compression. The constants q_1 , q_2 , and q_3 were introduced by Tvergaard in order to provide a better match with numerical studies [16]. Aravas provides a detailed explanation of implementing this model in a static finite element algorithm for three-dimensional solid elements [21]. The procedure used here is essentially the same, with some modifications due to the different element formulation. The stress is then transformed to local coordinates, and the internal force vector is computed as described above. One thing to be noted in Eq. (35) is that if Φ is initially 0, the yielding criteria surface is identical to the von Mises yield condition.

After calculating the strain tensor using Eq. (23), any previous plastic strain is subtracted using

$$\left\{ \varepsilon^{e} \right\} = \left\{ \varepsilon^{total} \right\} - \left\{ \varepsilon^{p} \right\} \tag{36}$$

The stress is then calculated using Eq. (24) with the components of $\{\varepsilon^e\}$. Using these values, the hydrostatic stress, deviatoric stress and effective stress are calculated as follows:

$$p = -\frac{1}{3} \left(\sigma_{xx} + \sigma_{yy} + \sigma_{zz} \right) \tag{37}$$

$$\{s\} = \{\sigma\} + p\{\delta\} \tag{38}$$

$$q = \sqrt{\frac{3}{2} \left(s_1^2 + s_2^2 + s_3^2 + 2 \left(s_4^2 + s_5^2 + s_6^2 \right) \right)}$$
 (39)

where δ is the Kronecker delta function:

$$\{\delta\} = \{11100\}^T \tag{40}$$

At this point, F is calculated from Eq. (35). If F is greater than zero, indicating plastic flow, iteration is required to determine the new porosity and change in plastic strain. The predictor-corrector method used in Aravas [21] is also used here. Using the values of p and q previously calculated as a first guess, correction factors are calculated using

$$\{C_t\} = [\mathbf{A}]^{-1} \{\mathbf{A}1\} \tag{41}$$

where

$$\{AI\} = \begin{cases} -F \\ -\Delta \varepsilon_p \frac{\partial F}{\partial q} - \Delta \varepsilon_q \frac{\partial F}{\partial p} \end{cases}$$
 (42)

$$\left[\mathbf{A} \right] = \begin{bmatrix} \mathbf{K} \frac{\partial F}{\partial p} + \frac{\partial F}{\partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta \varepsilon_p} & -3\mathbf{G} \frac{\partial F}{\partial q} + \frac{\partial F}{\partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta \varepsilon_q} \\ \frac{\partial F}{\partial q} + \Delta \varepsilon_q \left[\mathbf{K} \frac{\partial^2 F}{\partial p^2} + \frac{\partial^2 F}{\partial a \partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta \varepsilon_p} \right] + \Delta \varepsilon_p \frac{\partial^2 F}{\partial a \partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta \varepsilon_p} & \frac{\partial F}{\partial p} + \Delta \varepsilon_p \left[-3\mathbf{G} \frac{\partial^2 F}{\partial q^2} + \frac{\partial^2 F}{\partial a \partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta \varepsilon_q} \right] + \Delta \varepsilon_q \frac{\partial^2 F}{\partial a \partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta \varepsilon_q}$$
 (43)

$$\left\{C_f\right\} = \left\{\begin{matrix} \alpha \\ \beta \end{matrix}\right\} \tag{44}$$

and

$$\frac{\partial F}{\partial q} = \frac{2q}{\sigma_0^2}$$

$$\frac{\partial^2 F}{\partial q^2} = \frac{2}{\sigma_0^2}$$

$$\frac{\partial F}{\partial p} = 2q_1 \Phi \left(\frac{-3q_2}{2\sigma_0}\right) \sinh \left(\frac{-3q_2p}{2\sigma_0}\right)$$

$$\frac{\partial^2 F}{\partial p^2} = 2q_1 \Phi \left(\frac{3q_2}{2\sigma_0}\right)^2 \cosh \left(\frac{-3q_2p}{2\sigma_0}\right)$$

$$\frac{\partial F}{\partial \sigma_0} = \frac{-q^2}{2\sigma_0^3} + 2q_1 \Phi \frac{3q_2p}{2\sigma_0^2} \sinh \left(\frac{-3q_2p}{2\sigma_0^2}\right)$$

$$\frac{\partial^2 F}{\partial p \partial \sigma_0} = -2q_1 \Phi p \left(\frac{3q_2}{2\sigma_0^2}\right)^2 \cosh \left(\frac{-3q_2p}{2\sigma_0^2}\right) + \frac{3q_2}{2\sigma_0^3} \sinh \left(\frac{-3q_2p}{2\sigma_0^2}\right)$$

$$\frac{\partial^2 F}{\partial q \partial \sigma_0} = \frac{-q}{\sigma_0^3}$$

$$\frac{\partial^2 F}{\partial q \partial \sigma_0} = \frac{-q}{\sigma_0^3}$$
(45)

The values for α and β are used to correct the change in strain caused by hydrostatic pressure and the change in strain caused by effective stress (See Eq. (46)), which are then used to determine the change in the plastic strain vector, as shown in Eq. (47).

$$\Delta \varepsilon_p^{new} = \Delta \varepsilon_p^{old} + \alpha \quad (\Delta \varepsilon_p^0 = 0)$$

$$\Delta \varepsilon_q^{new} = \Delta \varepsilon_q^{old} + \beta \quad (\Delta \varepsilon_q^0 = 0)$$
(46)

$$\left\{\Delta \varepsilon^{p}\right\} = \frac{1}{3} \Delta \varepsilon_{p} \left\{\delta\right\} + \Delta \varepsilon_{q} \left(\frac{3}{2q}\right) \left\{s\right\} \tag{47}$$

This change in plastic strain, $\{\Delta_{\mathcal{E}}^{p}\}$, is then added to the total plastic strain, and the new elastic strain is calculated using Eq. (36), and the stress vector is calculated again using Eq. (24). Next, the change in void content, or porosity, is calculated as:

$$\Delta \Phi = \Delta \Phi_{growth} + \Delta \Phi_{nucleation} \tag{48}$$

$$\Delta\Phi_{growth} = (1 - \Phi) \sum \varepsilon_{ii}^{p} \tag{49}$$

$$\Delta \Phi_{nucleation} = \frac{\Phi_N}{s_N \sqrt{2\pi}} \exp \left[-\frac{1}{2} \left(\frac{\varepsilon^p - \varepsilon_N}{s_N} \right)^2 \right] \Delta \varepsilon_p$$
 (50)

where Φ_N is the volume fraction of void nucleating particles and ϵ_N and s_N are the mean and standard deviation of a normal distribution of nucleation strain, as suggested by Chu and Needleman [15] and utilized by Aravas [21]. Then the effective plastic strain, and void content are updated with

$$\Delta \varepsilon^{p} = \frac{-p \, \Delta \, \varepsilon_{p} + q \, \Delta \, \varepsilon_{q}}{\left(I - \Phi\right) \, \sigma_{\theta}} \tag{51}$$

$$\varepsilon_{t+\Delta t}^{p} = \varepsilon_{t}^{p} + \Delta \varepsilon^{p} \tag{52}$$

where ε_i^p is the effective plastic strain from the previous time step. At this point, the change in yield stress due to strain hardening is calculated (see below). If either α or β is greater than a predetermined tolerance, the process iterates beginning with Eq. (41). The tolerance used for all examples presented in the paper is 1.0×10^{-4} .

B. IMPROVING STABILITY IN THE CONSTITUTIVE EQUATIONS

Stability in the procedure outlined above is very dependent on the order in which the various equations are evaluated. Aravas discusses the stability problem when considering problems involving large plastic strains [21]. While the change in plastic strain from one time step to the next will theoretically be small, high strain rate and changes in the strain-hardening characteristics of the material act to degrade stability. Equations (42), (43) and (45) are not complete in that they do not contain all of the derivative terms required by the chain rule.

The predictor-corrector method used is based on Newton's method, where the correction factors, α and β , are found from

$$\begin{bmatrix} f_{1,\Delta\varepsilon_p} & f_{1,\Delta\varepsilon_q} \\ f_{2,\Delta\varepsilon_p} & f_{2,\Delta\varepsilon_q} \end{bmatrix} \begin{pmatrix} \alpha \\ \beta \end{pmatrix} = \begin{cases} f_1 \\ f_2 \end{cases}$$
(53)

where

$$f_{1} = \left(\frac{q}{\sigma_{0}}\right)^{2} + 2q_{1}\Phi \cosh\left(\frac{-3q_{2}p}{2\sigma_{0}}\right) - \left(1 + q_{3}\Phi^{2}\right)$$
 (54)

$$f_2 = \Delta \varepsilon_p \frac{\partial f_1}{\partial q} + \Delta \varepsilon_q \frac{\partial f_1}{\partial p}$$
 (55)

$$f_{1,\Delta\varepsilon_p} = K \frac{\partial f_1}{\partial p} + \frac{\partial f_1}{\partial \Phi} \frac{\partial \Phi}{\partial \Delta\varepsilon_p} + \frac{\partial f_1}{\partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta\varepsilon_p}$$
 (56)

$$f_{1,\Delta\varepsilon_q} = -3G\frac{\partial f_1}{\partial q} + \frac{\partial f_1}{\partial \Phi} \frac{\partial \Phi}{\partial \Delta\varepsilon_q} + \frac{\partial f_1}{\partial \sigma_0} \frac{\partial \sigma_0}{\partial \Delta\varepsilon_q}$$
 (57)

$$f_{2,\Delta\varepsilon_{p}} = \frac{\partial f_{1}}{\partial q} + \Delta\varepsilon_{p} \frac{\partial^{2} f_{1}}{\partial \sigma_{0} \partial q} \frac{\partial \sigma_{0}}{\partial \Delta\varepsilon_{p}} + \Delta\varepsilon_{q} \left[K \frac{\partial^{2} f_{1}}{\partial p^{2}} + \frac{\partial^{2} f_{1}}{\partial p \partial \Phi} \frac{\partial \Phi}{\partial \Delta\varepsilon_{p}} + \frac{\partial^{2} f_{1}}{\partial p \partial \sigma_{0}} \frac{\partial \sigma_{0}}{\partial \Delta\varepsilon_{p}} \right]$$
(58)

$$f_{2,\Delta\varepsilon_{q}} = \frac{\partial f_{1}}{\partial p} + \Delta\varepsilon_{q} \left[\frac{\partial^{2} f_{1}}{\partial p \partial \Phi} \frac{\partial \Phi}{\partial \Delta\varepsilon_{q}} + \frac{\partial^{2} f_{1}}{\partial p \partial \sigma_{0}} \frac{\partial \sigma_{0}}{\partial \Delta\varepsilon_{q}} \right] + \Delta\varepsilon_{p} \left[-3G \frac{\partial^{2} f_{1}}{\partial q^{2}} + \frac{\partial^{2} f_{1}}{\partial q \partial \sigma_{0}} \frac{\partial \sigma_{0}}{\partial \Delta\varepsilon_{q}} \right]$$
(59)

Derivative terms that are zero have been dropped, and K and G are the bulk and shear moduli, respectively. f_1 is Gurson's void function, and f_2 is the flow rule, both of which are driven to zero. As void content increases, the number of iterations required to achieve convergence increases dramatically, and often diverges, instead. By removing all partial derivatives relating to void content, the stability of the algorithm is greatly increased, at the cost of only 2 to 3 extra iterations, depending on the current void content. The dependence

on the current yield stress is retained to allow convergence across a transition in the slope of the yield stress versus strain relationship.

The partial derivatives of yield stress with respect to $\Delta \varepsilon_{\rm p}$ and $\Delta \varepsilon_{\rm q}$ are computed by calculating the slope from the current yield stress to the yield stress corresponding to the increase in plastic strain from the corrected values of $\Delta \varepsilon_{\rm p}$ and $\Delta \varepsilon_{\rm q}$. This means that these derivatives will be zero on the first iteration, since both of the control variables are initialized to zero. Several error traps must be included to prevent round-off and truncation error from causing the algorithm to diverge, and to prevent porosity from decreasing or becoming negative. One of the assumptions used in this implementation is that once voids form, they do not disappear. In other words, voids do not disappear when the element is placed in compression.

C. STRAIN HARDENING

Modeling the nonlinear elasto-plastic behavior of the material used is simplified by constructing a piece-wise linear version of the stress-strain plot. Using the tangent modulus, E_{T} , for each piece-wise region, the yield stress is calculated with

$$\sigma_0^{t+\Delta t} = \sigma_0^0 + \sum_{i=1}^{j-1} E_{Ti} \left(\varepsilon_i - \varepsilon_{i-1} \right) + E_{Tj} \left(\varepsilon_i^{eff} - \varepsilon_{j-1} \right)$$
 (60)

where $\sigma_0^{t+\Delta t}$ is the yield stress for this time step, σ_0^0 is the original yield stress, ε_i^{eff} is the total effective strain for the time step under consideration, and ε_i is the upper strain limit of the *i*th linear segment. ε_0 is the original yield strain, and is calculated by the program as

simply σ_0^0/E . The current effective elastic strain is calculated by dividing the current yield stress by the elastic modulus, E. This is added to the current effective plastic strain to obtain the total effective strain.

As an example, let the total effective strain from Eq. (49) lie within the second work-hardening segment. The new yield stress is

$$\sigma_0^{t+\Delta t} = \sigma_0^0 + E_{TI} \left(\varepsilon_I - \varepsilon_0 \right) + E_{T2} \left(\varepsilon_{t+\Delta t}^{eff} - \varepsilon_1 \right) \tag{61}$$

Figure 3 illustrates this example. The algorithm calculates the new yield stress as a function of the cumulative effective plastic strain, the current effective elastic strain, and the original yield stress for each iteration of the damage constitutive equations.

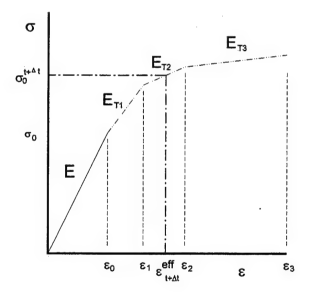


Figure 3. Calculating a New Yield Stress.

IV. HOURGLASS MODE CONTROL

Only one integration point is used in each plane parallel to the reference plane, which results in under-integration in the ξ - η plane. This leads to spurious, hourglass, or zero-energy modes in the element, which will yield useless results if left uncontrolled. The effects of these modes are shown in Fig. 4, a pinched cylinder where one eighth of the structure is modeled by utilizing symmetry boundary conditions. Clearly, no useful information can be obtained from the resulting solution.

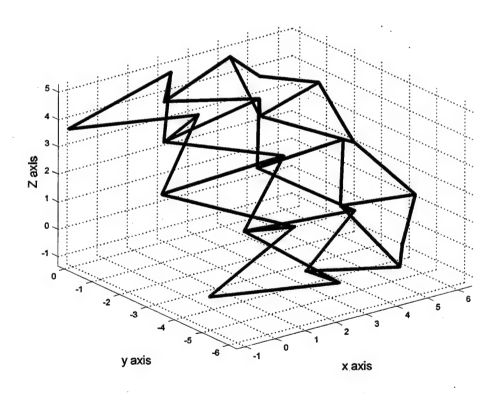


Figure 4. Hourglass Modes in a Pinched Cylinder Model.

Belytschko, et al., proposed an efficient means of controlling the hourglass modes of a similar element [25]. The method described uses a portion of a stiffness matrix generated by full integration in all directions to modify the stiffness matrix generated by under-integration. Although the formulation proposed in this paper does not use a stiffness matrix, a similar approach is just as effective in controlling these modes.

Rather than fully integrating in all three directions, the element is fully integrated in the ξ - η plane, but under-integrated in the ζ direction, as shown in Fig. 5. The procedure described above for calculating the internal force vector is followed to generate an internal force vector related to these four integration points.

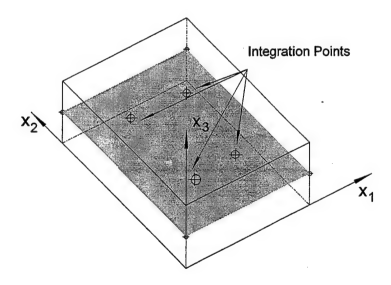


Figure 5. Hourglass Mode Control Integration Points.

The algorithm for calculating strain, stress, and force for the hourglass mode control integration points is identical to the algorithm used to produce the main internal force vector,

with the exception of plastic strain. The damage constitutive equations described in the previous section are not utilized for hourglass mode control. The internal forces generated from the two integration schemes are treated like the stiffness matrices in Belytschko et al. [25]. For nz integration points through the element thickness, the new force vector is

$$\left\{f_{int}\right\} = \left\{f_{int}^{IxIx\,nz}\right\} + h\left\{f_{hour}\right\} \tag{62}$$

where

$$\left\{ f_{hour} \right\} = \left\{ f_{int}^{2x2x1} \right\} - \left\{ f_{int}^{1x1xnz} \right\} \tag{63}$$

and

$$h = \frac{r t^2}{A} \tag{64}$$

The variable h is used here in Eqs. (62) and (64) instead of the ε used in Belytschko et al. [25], to avoid confusion with the various strains discussed in this paper. The variables used to calculate h are the element thickness (t) and the element's surface area (A). The effect of r follows that described in Belytschko et al. [25], and is set to 0.05. The range of values for r that effectively controls the hourglass modes, but does not greatly affect the overall element stiffness is roughly 0.046 to 0.057 (determined experimentally). Since the elements are in arbitrary orientation in 3-D space, the area calculation is computed as the sum of the area of the two triangles formed by dividing the element at the diagonal between nodes one and three:

$$A = \frac{1}{2} \sqrt{l_1^2 l_2^2 - D_1^2 + l_3^2 l_4^2 - D_2^2}$$
 (65)

where

$$D_{1} = (x_{1}^{1} - x_{1}^{2})(x_{1}^{3} - x_{1}^{2}) + (x_{2}^{1} - x_{2}^{2})(x_{2}^{3} - x_{2}^{2}) + (x_{3}^{1} - x_{3}^{2})(x_{3}^{3} - x_{3}^{2})$$

$$D_{2} = (x_{1}^{1} - x_{1}^{4})(x_{1}^{3} - x_{1}^{4}) + (x_{2}^{1} - x_{2}^{4})(x_{2}^{3} - x_{2}^{4}) + (x_{3}^{1} - x_{3}^{4})(x_{3}^{3} - x_{3}^{4})$$

$$(66)$$

and

$$l_{I} = \sqrt{\left(x_{1}^{2} - x_{1}^{1}\right)^{2} + \left(x_{2}^{2} - x_{2}^{1}\right)^{2} + \left(x_{3}^{2} - x_{3}^{1}\right)^{2}}$$

$$l_{2} = \sqrt{\left(x_{1}^{3} - x_{1}^{2}\right)^{2} + \left(x_{2}^{3} - x_{2}^{2}\right)^{2} + \left(x_{3}^{3} - x_{3}^{2}\right)^{2}}$$

$$l_{3} = \sqrt{\left(x_{1}^{4} - x_{1}^{3}\right)^{2} + \left(x_{2}^{4} - x_{2}^{3}\right)^{2} + \left(x_{3}^{4} - x_{3}^{3}\right)^{2}}$$

$$l_{4} = \sqrt{\left(x_{1}^{1} - x_{1}^{4}\right)^{2} + \left(x_{2}^{1} - x_{2}^{4}\right)^{2} + \left(x_{3}^{1} - x_{3}^{4}\right)^{2}}$$
(67)

and (x_1^k, x_2^k, x_3^k) is the location of node k in the global coordinate system. For these calculations, the element is assumed to be flat (no curvature along either the ξ or η directions). The effectiveness of this method of control is shown in Fig. 6, the same problem as shown in Fig. 4, but with the hourglass mode control described above.

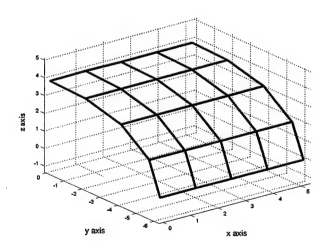


Figure 6. Pinched Cylinder Model with Hourglass Mode Control.

All verification problems analyzed in the following section were completed with hourglass control enabled. The current implementation uses hourglass control consistently, but allows easy modification to make hourglass control a user-defined option. The internal hourglass mode control can be disabled when the element is used in a general finite element program that includes various methods of spurious mode control.

V. IMPLEMENTATION IN RESEARCH-ORIENTED FE CODE

An in-house general purpose finite element analysis program, referred to herein as FEA, was used to verify the algorithms discussed. Several of these verification problems are presented in the next section. The program is written in FORTRAN, and is tailored for easy modification, rather than speed or ability to handle very large problems. The program reads the system parameters from a single input file, then writes the results to two ASCII text files: one contains the nodal coordinates and displacements for all time steps, the other contains the data associated with each element for all time steps (stress and strain tensors, effective plastic strain, void content, yield stress, etc.).

A. PREPROCESSOR

Since this program is locally generated, there is no preprocessor or postprocessor available. In order to facilitate running several problems, a preprocessor was written in MATLAB that generates the proper input file for several simple geometric shapes. The MATLAB script file will create a properly meshed flat plate, open box, cylinder, complete spherical cap, or a spherical cap with a hole in its top. The listing for the preprocessor is presented in Appendix A.

B. POSTPROCESSOR

A postprocessor, also written as a MATLAB script file, utilizes a graphical user interface (GUI) to create the required graphs. It also has the capability to create an animated

display of the displacement over time. The displacements are scaled to make them clearly visible. This feature is useful in determining the effectiveness of hourglass control and verifying the boundary conditions applied. However, the MATLAB movie is memory intensive and slow, and is therefore useful only as a qualitative verification of proper problem definition. All graphs of problem solutions, except as noted, were generated using this postprocessor. The full listing of all associated files is provided in Appendix B.

C. FEA SHELL ELEMENT SUBROUTINE DESCRIPTION

The FORTRAN source code for the FEA implementation of the element presented here, along with the proposed hourglass control and constitutive model, is listed without comment in Appendix C. Since the primary purpose of the FEA program is to assist in the development of finite element algorithms, this subroutine is written with readability, rather than computational efficiency, as a primary concern,. A brief outline of the subroutine is shown in Fig. 7. The FEA program treats each type of element (shell, beam, etc.) in a single block, and loops through each element, then each integration point within the element.

The order of calculation when evaluating the constitutive equations is critical, and any variation will cause instability. The two control variables, dep and deq, are updated by the correction factors on each iteration. All other variables are initialized to the values from the previous time step within each iteration. This essential feature allows stability of the algorithm. As discussed above, the variation of void content with dep and deq is not used,

but the variation of yield stress with the two control variables is calculated and included in the full expansion of the partial derivatives.

Program Outline for ELSH4L.F				
1. 2. 3. 4.	Enter subroutine Set constants Load gauss points and weights Begin loop over each shell element A. Initialize element internal force vectors B. Retrieve element material properties C. Compute element indices into system vectors D. Load nodal coordinates and displacements E. Compute local coordinates system and surface area F. Compute rotation matrices G. Transform displacements to local coordinate system H. Calculate hourglass force vector I. Begin Integration Loop 1) Load shape functions and derivatives 2) Calculate Jacobian, its inverse, and its determinate 3) Calculate strain-displacement matrix ([B]) 4) Compute strain ({e} = [B] {d}) 5) Transform strain to local coordinate system 6) Subtract any previous plastic strain from strain tensor 7) Load last values of σ_{c_0} ϕ_0 and e^{ρ}_{eff} 8) Compute stress ($\{\sigma\} = [D]\{\epsilon\}$) 9) Compute yield function (F) 10) If $F > 0$, set dep = deq = 0 and begin iteration a) Calculate required partial derivatives b) Prevent Numerical Errors (Divide by Zero, etc.) c) Retrieve Previous ϕ_0 , e^{ρ}_{eff} , and σ_0 d) Update dep and deq e) Compute Void Growth g) Compute New ϕ_{ij} , δ_{ij} , and e^{ρ}_{eff} h) Compute New ϕ_{ij} , δ_{ij} , and δ^{ρ}_{eff} h) Compute New ϕ_{ij} , δ_{ij} , and δ^{ρ}_{eff} i) Compute New ϕ_{ij} , δ_{ij} , and δ^{ρ}_{eff} h) Transform stress to global coordinate system 10) Last values used are retained. 12) Transform stress to global coordinate system 13) Calculate internal force for this integration point and sum 14) Store material tensors and constants in system vectors 15) End of integration loop J. Apply hourglass correction force K. Transform iternal force vector to global coordinate system L. Compute element mass vector M. Load mass and force vectors into system vector			

Figure 7. Outline of FEA Implementation.

VI. NUMERICAL EXAMPLES

The element presented here has been extensively tested using a variety of problems in which an analytic solution was available, and has produced satisfactory results in all cases studied. The transformation matrices and constitutive equations were verified with specifically tailored problems, and the element passes the patch test. The examples presented below are representative of the test cases used to verify the element, and are presented in the order of curvature: flat, singly curved, and doubly curved. For each case, an elastic solution is compared to available results, and then the elasto-plastic solutions are presented.

A. ELASTIC PLATE

A plate clamped on all four sides is subjected to a concentrated force at its center. The elastic modulus is 10 Msi (68.95 GPa), the density is 0.1647 slugs/in³ (0.147 kg/cm³), and Poisson's ratio is 0.2. The dimensions of the plate are 10 in x 10 in x 0.1 in thick (25.4cm x 25.4 cm x 0.254 cm). The yield stress is set high enough to ensure a completely elastic response. Two integration points through the thickness are used. The applied force is 40 lbf (177.9 N). The finite element mesh uses symmetry to model one quarter of the plate, with appropriate symmetry boundary conditions applied. Both a 2x2 (4 element) and 4x4 (16 element) mesh are used in the finite element analysis. The results for both meshes and the analytic solution are shown in Table 1. Using a four-element mesh, the new shell

element obtained a displacement within 3.27% of the analytic solution, and 0.82% using a 16 element mesh.

Table 1. Comparison of Results for Elastic Clamped Plate.

Analysis Type	Center Node Peak Displacement (in)
2 x 2 FE Mesh (Dynamic)	-4.74x10 ⁻²
4 x 4 FE Mesh (Dynamic)	-4.94x10 ⁻²
Analytic (Twice the Static Solution)	-4.90x10 ⁻²

B. THICK CLAMPED PLATE UNDER PRESSURE LOAD IN ELASTO-PLASTIC REGION

A thick steel plate, clamped on all four sides, is subjected to dynamic pressure load.

The plate is 6m by 6m by 0.6m thick (for a 10:1 ratio). Table 2 shows the material properties for the structure. Table 3 shows the properties for the void model.

Table 2. Material Properties of Clamped Plate.

Property	Value	Units
Elastic Modulus (E)	2x10 ¹¹	Pa
Tangent Modulus (E _r)	2x10 ¹⁰	Pa
Density (ρ)	7850	kg/m³
Poisson's ratio (v)	0.29	(none)
Yield Stress (Syp)	2.5x10 ⁸	Pa

Table 3. Void Characteristics of Clamped Plate.

Initial Void Content (Φ_0)	0.0	
Nucleating Particle Content (Φ _N)	0.04	
Mean Nucleation Strain (e _N)	0.3	
Nucleation Strain Standard Deviation (s _N)	0.1	
Model Constant q ₁	1.5	
Model Constant q ₂	1.0	
Model Constant $q_3 (= q_1^2)$	2.25	

One quarter of the plate is modeled with a three by three element mesh, for a total of nine elements, with appropriate symmetry boundary conditions applied, and is shown below in Fig. 8.

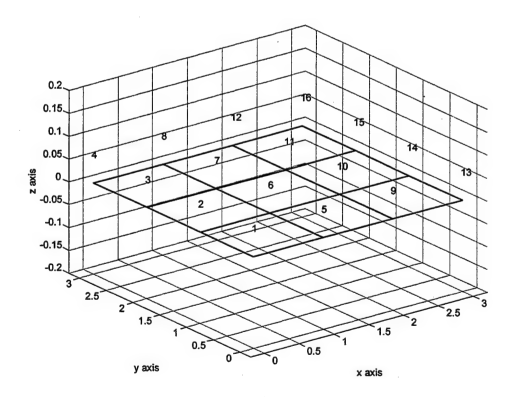


Figure 8. Nine Element Clamped Plate Mesh.

Four integration points are used through the thickness of each element. The calculation time step is 10^{-5} seconds, while the data is plotted at 2×10^{-4} second increments. The analysis terminates at 0.04 seconds. The plate is subjected to a uniformly distributed pressure acting downwards, and includes the appropriate drilling moment. The pressure increases linearly from 0.0 Pa at the start to 80 MPa at 0.01 seconds, then remains constant for the duration of the analysis. Three cases were analyzed: no void effects, void growth effects only, and void growth and nucleation. Figure 9 illustrates the effect of voids on the effective stress versus the effective strain relationship in the top of one of the border elements. When void effects are not included, the Von-Mises Equivalent (VME) stress follows the yield stress in the plastic region (See Fig. 9a).

Including void effects causes the yielding before the VME stress reaches the yield stress. As the void content increases with continued plastic flow, the VME stress falls farther below the yield stress, as shown in Fig. 9b (see Eq. (35)). Table 4 summarizes the results of the three cases. The most significant difference is in the transverse normal stress, σ_{zz} . This difference resulted in a 2.0% increase in the effective plastic strain, and a 2.6% increase in the peak deflection of the center of the plate.

Another interesting point is that the transverse normal stress was compressive and constant throughout the thickness up until plastic flow, as assumed in the formulation, then varied as the stress in the bottom fiber decreased and became tensile. Figure 10 shows the variation of transverse normal stress through the thickness of the element at the time of peak stress. Eventually, the transverse normal stress varied from compressive at the top, where

the pressure was applied, and tensile at the bottom. The examples that follow will not include a separate analysis for void growth effects only, since the difference of including nucleation effects at the resulting small plastic strains obtained do not cause significantly different results.

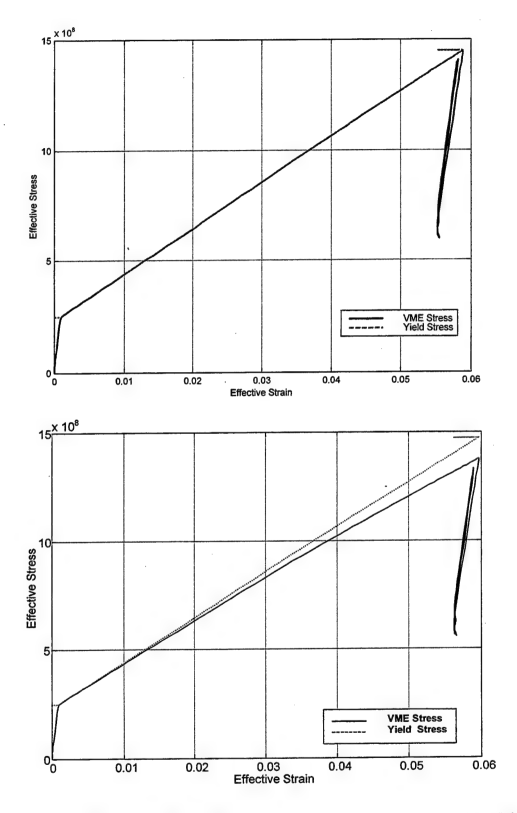


Figure 9. Void Effects in a Clamped Plate with Pressure Loading: Top (a) - No Void Effects, Bottom (b) - Void Effects.

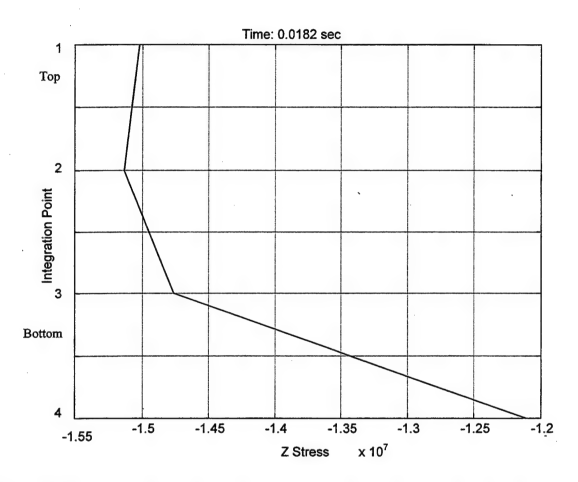


Figure 10. Transverse Normal Stress Variation through Shell Thickness: Clamped Plate with Pressure Loading and Void Effects.

Table 4. Summary of Results for Clamped Plate with Pressure Load.

Peak Values for Element #3	No Void Effects	Void Growth	Growth and Nucleation
σ _{VM} (Gpa)	1.4498	1.3789	1.3784
€ _{effective}	0.0588	0.0597	0.0597
σ _{xx} (GPa)	1.4501	1.3603	1.3598
σ _{yy} (GPa)	0.6561	0.6030	0.6073
σ _{zz} (GPa) (Max Tension)	-0.0003	0.0065	0.0091
σ _{zz} (GPa) (Max Compression)	-0.0363	-0.0322	-0.0320
ε ^{plastic} (effective)	0.0540	0.0551	0.0551
Φ (Porosity)	NA	0.0355	0.0357
Deflection (m) (Center Node)	-0.3801	-0.3897	-0.3898

C. THICK CLAMPED PLATE WITH CENTRAL POINT LOAD IN THE ELASTO-PLASTIC REGION

The geometry and material properties of this example are identical to those used in the previous example. The pressure load has been replaced with a point load at the center node. Four cases are studied: without void effects or a drilling moment, with void effects, with a drilling moment applied, and with both void effects and a drilling moment applied. All four sides are clamped, and the center node displacement is restricted in the x and y directions. Then, a DM equal to the applied force times one-half the plate thickness is applied for both the no-void and void cases. The time history of the stress components in the center element is shown in Fig. 11, and the relationship between porosity and effective plastic strain is illustrated in Fig. 12. The results for all six cases are summarized below in Table 5.

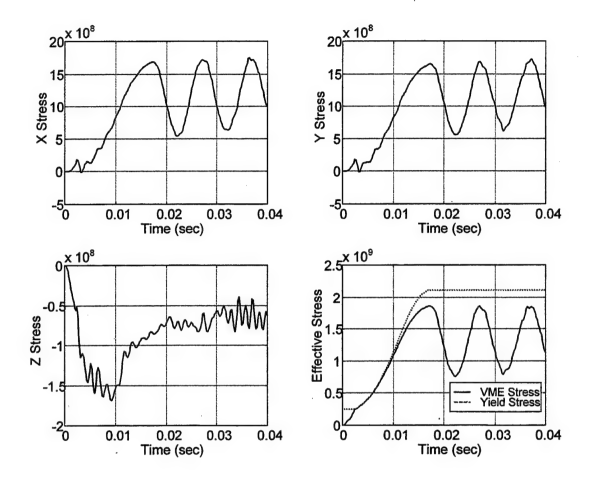


Figure 11. Stress Component Time History in Bottom Fiber: Clamped Plate with Concentrated Load, Void Effects, and Drilling Moment Applied.

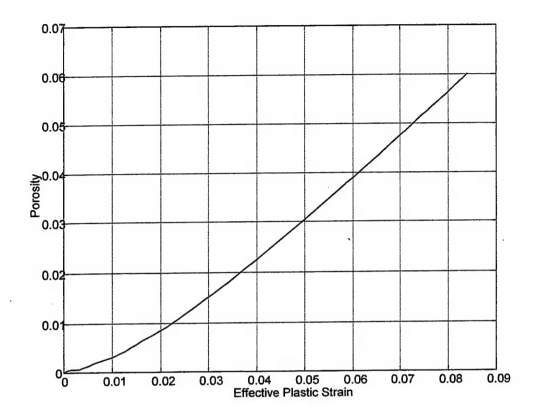


Figure 12. Porosity versus Effective Plastic Strain in Bottom Fiber: Clamped Plate with Concentrated Load, Void Effects, and Drilling Moment Applied.

Table 5. Summary of Results for Clamped Plate Subjected to Point Force.

Peak Values for Center	No Voids	Voids	No Voids	Voids
Element	No DM	No DM	Drilling Moment	Drilling Moment
σ _{vm} (GPa)	2.1119	1.9021	2.0540	1.8590
E _{effective}	0.0892	0.0929	0.0865	0.0887
σ _{xx} (GPa)	2.0008	1.8407	2.0000	1.7498
σ _{vv} (GPa)	1.9588	1.7245	1.8589	1.7234
σ _{zz} (GPa) (Max Tension)	0.0048	0.0213	-0.0003	-0.0003
σ _{zz} (GPa) (Max Compression)	-0.0036	-0.0057	-0.1751	-0.1691
ε ^{plastic} (effective)	0.0838	0.0880	0.0812	0.0840
Φ (Porosity)	NA	0.0663	NA	0.0601
Deflection (m) (Center Node)	-0.4413	-0.4544	-0.4350	-0.4451

The drilling moment increases the effective stress on the fiber in compression, and decreases the effective stress on the fiber in tension. The reduction in the tensile stress results in a reduction in the effective plastic strain. Including void effects decreases the VME stress on the fiber in tension, and slightly increases the VME stress on the fiber in compression. The effective plastic strain is also increased. All of these results indicate that this formulation correctly predicts, in a qualitative sense, the effects of drilling moments and void growth and nucleation. Including only void effects increased the effective plastic strain by 5.0%, while including only the drilling moment decreased the effective plastic strain by 3.1%. Including both void effects and the drilling moment increased the effective plastic strain by 0.2%, indicating the importance of applying both effects together.

D. SIMPLY SUPPORTED PLATE WITH CENTRAL POINT LOAD IN THE ELASTO-PLASTIC REGION

The material properties, geometry, mesh, and time values from the clamped plate problem above are used here. The magnitude of the force is $8x10^8$ N, and the drilling moment, when applied, is $-2.4x10^8$ N. The same four cases are analyzed: no void or drilling moment effects, drilling moment effects only, void effects only, and both void and drilling moment effects. The analysis results for all four cases are summarized in Table 6.

Table 6. Summary of Results for a Simply Supported Plate with a Point Force.

Peak Values for Center	No Voids	No Voids	Voids	Voids and Drilling Moment
Element	No DM	Drilling Moment	No DM	
σ _{VM} (GPa)	3.0874	3.0089	2.5061	2.4881
E _{effective}	0.1360	0.1321	0.1450	0.1393
σ _{xx} (GPa)	3.0253	2.9123	2.4686	2.4206
σ _{vv} (GPa)	3.0459	2.8889	2.4612	2.3928
σ _{zz} (GPa) (Max Tension)	0.0060	-0.0003	0.0114	-0.0003
σ _{zz} (GPa) (Max Compression)	-0.0035	-0.1802	-0.0035	-0.1722
ε ^{plastic} (effective)	0.1277	0.1242	0.1382	0.1327
Φ (Porosity)	NA	NA	0.1197	0.1092
Deflection (m) (Center Node)	-0.9219	-0.8990	-0.9770	-0.9420

The qualitative results for this example are the same as for the clamped plate. Since the applied force causes greater initial yielding, void growth and nucleation has a greater effect. The drilling moment, when added to the void effects, reduces the amount of effective plastic strain in tension and displacement.

E. ELASTIC PINCHED CYLINDER

An open-ended cylinder of radius 5.0 in. (12.7 cm), length 10.35 in. (26.289 cm), and thickness 0.094 in. (0.23876 cm) is subjected to a pinching load of 100 lbf (444.82 N) (See Fig. 6). The elastic modulus is 10.5 msi (72.4 Gpa), Poisson's ratio is 0.3125, and the density is 3.125x10⁻³ slugs/in³ (2783 kg/m³). The load is applied as a step function beginning at time t = 0 seconds. Using symmetry, the problem was reduced to a one-eighth section of the cylinder. The dynamic value should be twice the analytic static value. Inextensional shell theory gives a static radial contraction of 0.1117 in. (0.28372 cm). The maximum radial contraction of the model is 0.1995 in. (0.5067 cm), which translates to a static radial contraction of 0.09975 in (0.2534 cm). Using a 256-element mesh (16 by 16), the maximum radial contraction was 0.2207 in. (0.5606 cm), for a static contraction of 0.1104 in (0.2804 cm). The results are summarized in Table 7. The 16-element solution is within 10.7% of the analytic solution, while the 256-element mesh is within 1.21%.

Table 7. Comparison of Results for Elastic Pinched Cylinder.

Analysis Type	Radial Contraction (in)
Finite Element with 16 Element Mesh	0.1995 (0.5067 cm)
Finite Element with 256 Element Mesh	0.2207 (0.5606 cm)
Analytic (Twice the Static Solution)	0.2234 (0.5674 cm)

F. THICK PINCHED CYLINDER IN THE ELASTO-PLASTIC REGION

The material and void properties shown in Tables 3 and 4 are used for an open-ended cylinder with a pinching force at its center. The top half of the cylinder is modeled using a 36 element mesh with appropriate symmetry boundary conditions along the bottom edge of mesh, as shown in Fig. 13. The analysis is calculated in 10⁻⁵ second steps, with output every $2x10^{-4}$ seconds, from 0.0 seconds to 0.04 seconds. The pinching force is 66.792 kN on each side, with a drilling moment of 848.258 Nm applied on the cases indicated. The same four cases are compared: no voids or drilling moment, void effects only, drilling moment only, and both void effects and drilling moment (DM). The effects of void growth and nucleation and the drilling moment is shown in the stress component time histories; Fig. 14 shows the stresses without voids or a drilling moment, and Fig. 15 shows the stresses with both effects included. The results for all cases are summarized in Table 8.

Table 8. Summary of Results for Elasto-Plastic Pinched Cylinder.

Peak Values for Center	No Voids	No Voids	Voids	Voids and
Element	No DM	Drilling Moment	No DM	Drilling Moment
σ _{VM} (MPa)	262.40	265.05	259.28	283.49
ε _{effective} (x 10 ³)	1.6504	1.5722	1.4273	2.7049
σ _{xx} (MPa)	182.46	160.38	170.12	150.31
σ _{vv} (MPa)	291.71	283.77	291.83	297.97
σ _{zz} (MPa) (Max Tension)	5.0619	-0.0238	1.0270	7.5235
σ _{zz} (MPa) (Max Compression)	-7.6358	-28.739	-7.0364	-35.373
ε ^{plastic} (effective) (x 10 ³)	0.7828	0.7583	0.5771	1.6923
Φ (Porosity) (x 10 ³)	NA	NA	0.4773	1.1970
Deflection (mm) (Center Node) Global Maximum	2.7589	12.919	-0.0220	-0.0214
Deflection (mm) (Center Node) Global Minimum	-31.750	-31.750	-31.750	-31.750

Since there is only a small amount of plastic strain, the effects of void growth and nucleation are greatly reduced. Even with this small amount of plastic flow, the effective plastic strain was increased by over 116 percent with the inclusion of both drilling moment and void effects, while the peak center node displacement was only increased by approximately 4.5 percent.

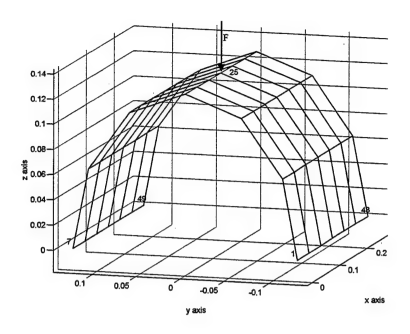


Figure 13. Mesh Structure for Pinched Cylinder.

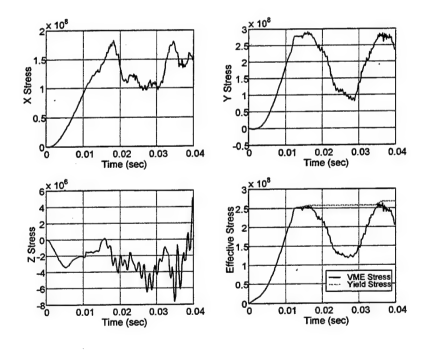


Figure 14. Stress Component Time History in Bottom Fiber: Pinched Cylinder, No Void Effects or Drilling Moment.

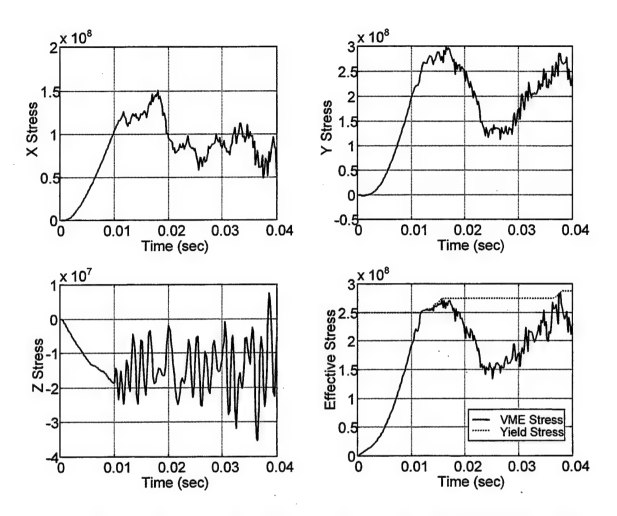


Figure 15. Stress Component Time History in Bottom Fiber: Pinched Cylinder, Void Effects and Drilling Moment Applied.

G. ELASTIC SPHERICAL CAP WITH A CENTER HOLE

The results of analysis with the shell element developed here are compared to results of the problem proposed in MacNeal and Harder [26]. The structure is a hemisphere of radius 10 units with a thickness of 0.04 units, and has an 18° hole cut in the center. Taking

advantage of the symmetry in the structure, only $\frac{1}{4}$ of the structure is modeled. An eight-by-eight mesh is used, for a total of 64 elements, with four integration points through the thickness of each element. The mesh used is shown in Fig. 16. The material properties of the structure are: $E = 6.825 \times 10^7$, $\rho = 0.001$, and $\nu = 0.3$. No void or drilling moment effects are employed in this example. Opposing forces of magnitude 2.0 are applied at each quadrant: the force at node 73 is of magnitude -1.0 and parallel to the y-axis, and the force at node one is of magnitude 1.0 and parallel to the x-axis. To obtain a representative static response using a dynamic model, the applied force begins with 0.0 magnitude at time 0, and increases linearly with a rise time of 0.16 seconds to the specified value. A calculation time step of 1×10^{-6} seconds is used, with termination at 0.22 seconds. The maximum deflection of node one was 0.0929, where the theoretical value is 0.0940, for a normalized displacement of 0.988. This is within the range of the results listed in MacNeal and Harder from the QUAD2 and QUAD4 elements, which are considered to be accurate for this problem [26]. These results are summarized in Table 9.

Table 9. Comparison of Results for Spherical Cap with Elastic Loading, and Results from MacNeal and Harder [26].

	New Shell Element	QUAD2	QUAD4
Normalized Displacement	0.988	0.986	1.005

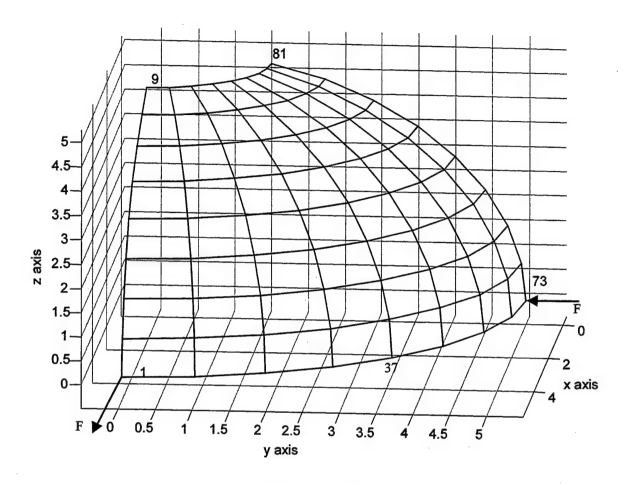


Figure 16. Mesh Structure for Spherical Cap.

H. SPHERICAL CAP WITH A CENTER HOLE, IMPACT LOADING

Key and Hoff [11] used the previous problem to test their element with an impact (step) load. The structure and matrial properties are the same, and the mesh is the same as shown in Fig. 16. The load is applied at its maximum at t=0.0⁺ seconds, rather than ramped up. Figure 17, which plots the displacement of node one obtained from both the shell element presented here and the element developed by Key and Hoff [11], shows that the

results of this element are comparable to those obtained by other elements under impact loading.

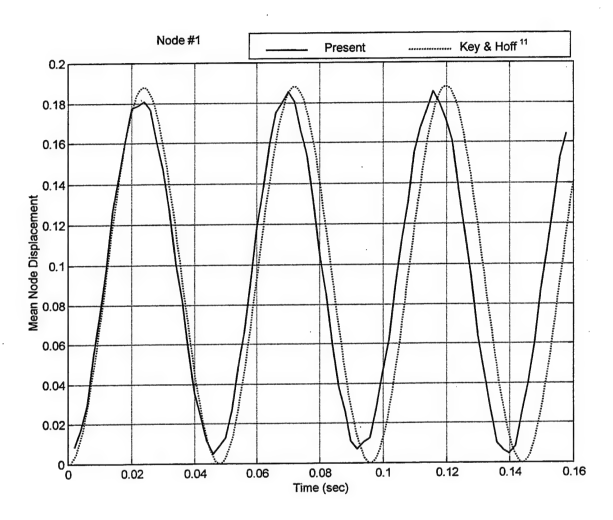


Figure 17. Node One Displacement Time History: Pinched Spherical Cap with Impact Loading.

I. SPHERICAL CAP WITH A CENTER HOLE, ELASTO-PLASTIC LOADING

The spherical cap structure shown in Fig. 16 is now used to verify the stability and convergence of the damage-constitutive equations under double curvature, and to illustrate

both the effects of void growth and nucleation and a drilling moment on a more complex structure. The material properties used are the same as shown in Tables 3 and 4. The structure has a radius of 5.0 meters and a thickness of 25 cm, for a radius to thickness ratio of 20:1, commonly considered the limit of thin-shell theory. Although a force is applied at each quadrant, all four loads are directed inwards. Therefore, only one load is applied at node 37 of the mesh shown in Fig. 16. The applied load is 5.9397x10⁶ N, with a drilling moment of -7.4246x10⁵ Nm applied for the cases indicated. A calculation time step of 1×10^{-5} seconds is used, with output every 5×10^{-4} seconds and stopping at 0.2 seconds. The load is initial zero, and increases linearly to its maximum at 0.01 seconds. The results shown in Table 10 are for the element nearest the applied load, where stress is maximum. The porosity versus effective plastic strain relationship is shown in Fig. 18. Due to the small amount of plastic strain, porosity is restricted to the linear zone, as shown. Figures 19a and 19b illustrate the effective stress versus effective strain in the same element on the compressive side and tensile side, respectively. Note that as the structure returns from a peak displacement, the stress follows the elastic modulus, not the tangent modulus. This reflects the correct behavior of material: the elastic modulus is not affected by plastic deformation, only the yield stress is affected. The node displacement where the force is applied is shown in Fig. 20. This figure shows that there are at least two vibration modes in operation.

Table 10. Summary of Results for Elasto-Plastic Spherical Cap.

Peak Values for	No Voids	No Voids	Voids	Voids and
Element #25	No DM	Drilling Moment	No DM	Drilling Moment
σ _{VM} (MPa)	250.26	250.76	250.26	250.75
$\varepsilon_{\text{effective}} (x \ 10^3)$	1.0730	1.1021	1.0731	1.1021
σ _{xx} (MPa)	202.09	199.44	202.08	199.43
σ _w (MPa)	-47.093	-58.183	-47.087	-58.192
σ _{zz} (MPa)	-5.2570	-12.433	-5.2571	-12.433
ε ^{plastic} (x 10 ⁵)	2.1002	4.1795	2.1020	4.1849
Φ (Porosity) (x 10 ⁶)	NA	NA	6.7149	12.763
Deflection (m) (Point of Loading)	0.0233	0.0246	0.0233	0.0246

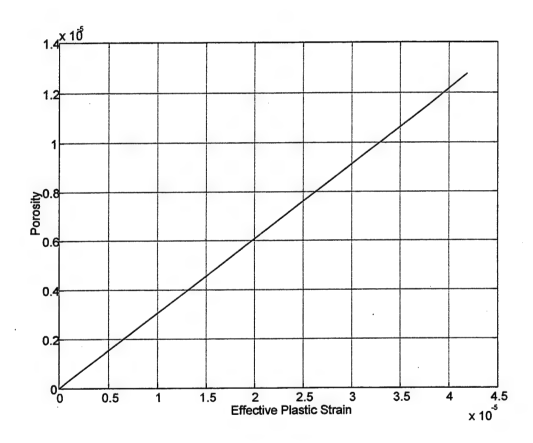


Figure 18. Porosity versus Effective Plastic Strain in Inner Fiber: Spherical Cap with Void and Drilling Moment Effects.

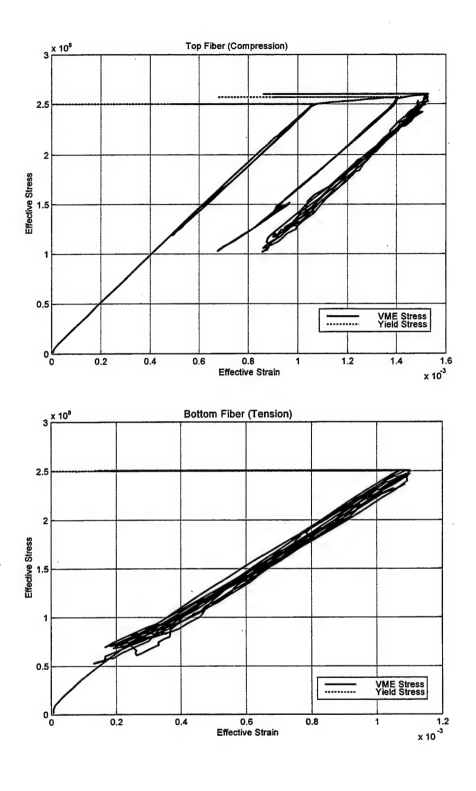


Figure 19. Comparison of Void and Drilling Moment Effects in a) Compression (Top) and b) Tension (Bottom) for Spherical Cap.

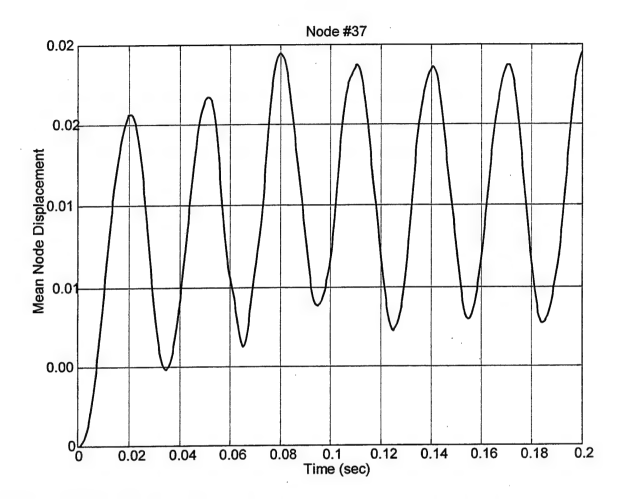


Figure 20. Contact Node Mean Displacement Time History: Spherical Cap with Void and Drilling Moment Effects.

During testing, it became apparent that this problem is not suitable for testing elastoplastic analysis. Since the structure is concave, the entire structure quickly collapses once yielding begins. In addition, the single point used to prevent rigid-body motion also provided a stress concentration and additional yielding (the "corner" began folding over). This necessitated using a force that just causes plastic flow, but does not collapse the structure or cause yielding near the anchored node. This is illustrated by an analysis of a 5 percent greater load than used for the analysis shown in Fig. 20. The resulting node 37 displacement is shown in Fig. 21, and the deformed structure in Fig. 22. However, it is still apparent that the qualitative results obtained in the previous elasto-plastic examples carried through to this problem; both voids and the drilling moment decreased stress in fibers under tension, and increased stress in fibers under compression. Due to the small amount of plastic strain, the increase in porosity was extremely small, which minimized the effects of the voids.

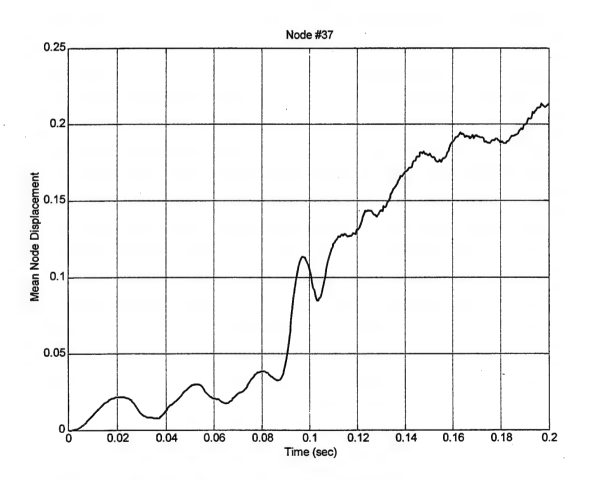


Figure 21. Contact Node Displacement: Spherical Cap with 5% Greater Load.

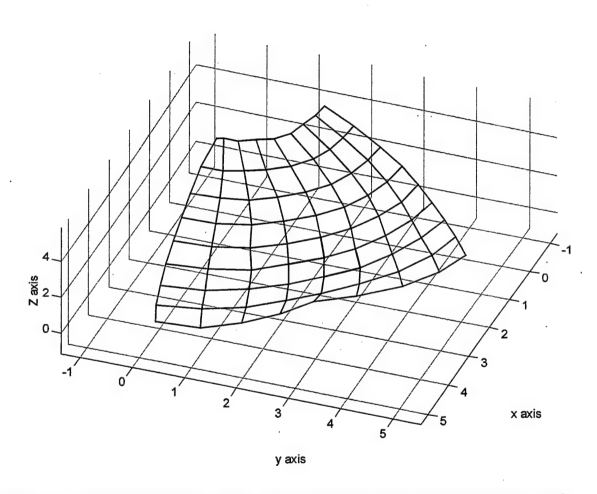


Figure 22. Deformed Structure at End of Analysis: Spherical Cap with 5% Greater Load.

VII. CONCLUSIONS AND RECOMMENDATIONS

A new shell formulation was developed for transient dynamic analysis which includes both void effects with plastic deformation and the transverse normal stress with drilling moment. The element is compatible with most shell elements which have three translation and three rotation degrees of freedom per node.

A numerically stable scheme was developed for Gurson's nonlinear constitutive equation model. The model includes both void nucleation and void growth. Furthermore, hourglass control was implemented into the algorithm to avoid spurious modes caused by the under-integration scheme.

The drilling moment was important for thick plates and shells. It induced large transverse normal stress and affected the plastic deformation. Similarly, the effect of voids on plastic deformation was not negligible. Numerical studies show that the effective plastic strain increased up to fifteen percent due to the combined effects of voids and the transverse normal stress.

The shell element presented here was tested on many cases, some of which had reference solutions for comparison. The new element gave accurate results to those problems.

Although the element formulation presented here performed well in numerical studies, the problems studied are not in the area where this element is most likely to be useful. The number of published problems available for comparison is limited, especially in the case of dynamic plastic deformation. The solutions provided by this element in the plastic region need to be verified by experimental data, and the constants of Gurson's void

model need to be determined for a variety of materials and porosity conditions. In all, there are six material property constants that must be determined before an accurate solution can be obtained: q_1 , q_2 , q_3 , f_N , e_N , and s_N . These are not readily available, but play a vital role in the constitutive equations of this formulation.

Implementing this shell element into a general purpose finite element program will allow easier comparisons with other element formulations, including solid brick elements, and will facilitate the analysis of more complex problems. This work has already begun for a version of DYNA3D, but is not yet complete.

The method of hourglass mode control used implemented a full version of the same formulation used for the internal force calculation. Efficiency may be increased by using a simpler formulation to calculate the internal hourglass forces. Along the same lines, no attempt has been made to improve the efficiency of the code presented here so that readability may be preserved.

APPENDIX A. PREPROCESSOR IN MATLAB FOR FEA.

```
% feapre.m - Preprocessor for fea program
      Program Limitations:
            No input of Initial Conditions
용
            Boundary Conditions must be applied manually to result file
            Load must be applied manually to result file
            Structure will be homogenous
            Only 1/4 cylinder supported at this time
% Get Input data
type = input('Type of Structure (1=cylinder, 2=plate, 3=shoe box,
4=cap1, 5=cap2):');
% Structure Specific Input
if type==1
      nelx = input('Number of Elements along x-axis:');
      nelp = input('Number of Elements perpendicular to x-axis:');
      x1 = input('Lower X-axis value:');
      x2 = input('Upper X-axis value:');
  R = input('Radius:');
   thetal = input('Starting Angle (degrees):');
  theta2 = input('Ending Angle (degress):');
elseif type==2
      nelx = input('Number of Elements along x-axis:');
      nelp = input('Number of Elements along y-axis:');
      x1 = input('Lower X-axis value:');
      x2 = input('Upper X-axis value:');
      y1 = input('Lower Y-axis value:');
      y2 = input('Upper Y-axis value:');
elseif type==3
      nelx = input('Number of Elements along x-axis:');
      nelp = input('Number of Elements Perpendicular to x-axis:');
      x1 = input('Lower X-axis value:');
      x2 = input('Upper X-axis value:');
      y1 = input('Lower Y-axis value:');
      y2 = input('Upper Y-axis value:');
      z1 = input('Lower Z-axis value:');
   z2 = input('Upper Z-axis value:');
elseif type==4
  R = input('Radius:');
  alpha = input('Phi (angle from Z-axis of inscribed circle):');
  ncap = input('Size (1=25 elem 2=42 elem 3=63 elem):');
  if ncap==1
     narc=6;
      ntheta=11;
   elseif ncap==2
     narc=8;
      ntheta=13;
   elseif ncap==3
      narc=10;
      ntheta=15;
   else
      errordlg('Bad Type');
   end
```

```
elseif type==5
   R = input('Radius:');
   alpha1 = input('Alpha1 (elevation angle to bottom ring (in
degrees):');
   alpha2 = input('Alpha2 (elevation angle to top ring (in degrees)):');
   thetal = input('Thetal (start angle from x-axis (in degrees)):');
   theta2 = input('Theta2 (end angle from x-axis (in degrees)):');
   nelp = input('Number of Elements along elevation angle:');
   nelx = input('Number of Elements along theta angle:');
else
      errordlg('Bad Type');
end
% Input Material and Time Data
E = input('Elastic Modulus:');
rho = input('Density (mass / unit volume):');
pois = input('Poissons Ratio:');
t = input('Thickness:');
Syp = input('Yield Stress:');
ipt = input('New(2) or Old(0) shell element (enter 2 or 0):');
stime = input('Start Time:');
etime = input('Stop Time:');
delt = input('Calculation Step Time (delta):');
ptime = input('Print Step Time:');
% If old shell element, adjust density to mass / unit area
if ipt==0
   rho = rho * t;
   ipoint = 0;
   ipoint = input('Number of Integration Points in Z direction:');
end
% *** Cylinder ***
if type==1
   % Calculate Node x, y, z coordinates
   thetal = thetal * pi / 180;
   theta2 = theta2 * pi / 180;
      nndsx = nelx+1;
      nndsr = nelp+1;
      xstep = (x2 - x1) / nelx;
      rstep = (theta2 - theta1) / nelp;
      i = 1;
      ne = nelx * nelp;
      nnodes = nndsx * nndsr;
      % Initialize Vectors
      x = zeros(nnodes, 1);
      y = x;
      z = x;
      nodes = zeros(ne,4);
      for xt=x1:xstep:x2
            for r=theta1:rstep:theta2
                  x(i) = xt;
                  y(i) = -R * cos(r);
                  z(i) = R * sin(r);
                  i = i + 1;
      end
   end
```

```
xd = x / R;
   yd = y / R;
   zd = z / R;
end
% *** Plate ***
if type==2
      nndsx = nelx+1;
      nndsy = nelp+1;
xstep = (x2 - x1)/nelx;
      ystep = (y2 - y1)/nelp;
      i = 1;
      ne = nelx * nelp;
      nnodes = nndsx * nndsy;
      x = zeros(nnodes, 1);
      y = x;
      z = x;
   nodes = zeros(ne, 4);
   xd = zeros(nnodes,1);
   yd = xd;
   zd = -1 * ones(nnodes, 1);
      for xt = x1:xstep:x2
             for yt=y1:ystep:y2
                   x(i) = xt;
                    y(i) = yt;
                    i = i + 1;
             end
      end
end
% *** Shoe Box ***
if type==3
   nndsx = nelx + 1;
      nndsp = nelp + 1;
      xstep = (x2 - x1)/nelx;
      ystep = (y2 - y1)/(nelp / 2);
zstep = (z2 - z1)/(nelp / 4);
      i = 1;
      ne = nelx * nelp ;
      nnodes = nndsx * nndsp;
      x = zeros(nnodes, 1);
      y = x;
   z = x;
   xd = x;
   yd = x;
   zd = x;
      nodes = zeros(ne,4);
      for xt = x1:xstep:x2
             % Front
             for zt = z1:zstep:z2
                   x(i) = xt;
          y(i) = y1;
          yd(i) = 1;
                    z(i) = zt;
                    i = i + 1;
             end
```

```
qoT &
            for yt = y1+ystep:ystep:y2
                  x(i) = xt;
                  y(i) = yt;
         z(i) = z2;
         zd(i) = -1;
                  i = i + 1;
            end
            % Back
            for zt = z2-zstep:-zstep:z1
                  x(i) = xt;
         y(i) = y2;
         yd(i) = -1;
                  z(i) = zt;
                  i = i + 1;
            end
      end '
end
% Spherical Cap #1
if type==4
   % Generates FEA mesh of spherical cap using symmetry
      % Also handles nodal connectivity
      % Input Values
  theta = linspace(0,pi/2,ntheta);
   alpha = pi * alpha / 180;
      phi = linspace(0,alpha,narc);
      % Spherical Coordinates for bottom nodes
      for i=1:ntheta
      P(i,:) = [R theta(i) phi(narc)];
      bct(i) = 7;
      bcr(i) = 7;
      end
      % Spherical Coordinates for next row of nodes
      nnodes = ntheta;
      for i = 1: (ntheta-1)/2
            P(i+ntheta,:) = [R theta(2 * i) (phi(narc)-(phi(narc)-...
                              phi(narc-1))/2)];
      end
      % Spherical Coordinates for the rest of the nodes
      nnodes = nnodes + (ntheta-1)/2;
     nrow = ntheta - (ntheta-1)/2;
      for j=narc-1:-1:1
      Th = 0;
      dTh = pi / (2 * (nrow - 1));
      bct(nnodes+1)=2;
      bcr(nnodes+1)=6;
      for i=1:nrow
            P(i+nnodes,:) = [R Th phi(j)];
            Th = Th + dTh;
         end
      nnodes = nnodes + nrow;
      bct(nnodes)=1;
      bcr(nnodes)=5;
         nrow = nrow - 1;
      end
```

```
% Change BC of top node
   bct(nnodes)=4;
   bcr(nnodes)=7;
       % Convert to Cartesian Coordinates
      for i=1:nnodes
          x(i) = P(i,1) * cos(P(i,2)) * sin(P(i,3));
          y(i) = P(i,1) * sin(P(i,2)) * sin(P(i,3));
          z(i) = P(i,1) * cos(P(i,3));
      end
      % Element Connections
      % Bottom row first
      mid = ntheta;
      top = mid + ntheta - (ntheta - 1)/2 -1;
      ne = 1;
      for i=1:(ntheta-1)/2,
          c1 = (i-1)*2 +1;
          nodes(ne,:) = [c1 (c1+1) (mid+i) (top+i)];
          nodes(ne+1,:) = [(c1+1) (c1+2) (top+1+i) (mid+i)];
          ne = ne + 2;
      % Rest of elements
      nrow = ntheta - (ntheta-1)/2 - 1;
      bot = mid;
      mid = top;
      top = top + nrow + 1;
      for j=1:nrow
          for i=1:nrow
             c1 = mid + i;
             nodes(ne,:) = [c1 (bot+i) (c1+1) (top+i)];
            ne=ne+1;
      end
          bot=mid+1;
          mid=top;
          top=top+nrow;
          nrow=nrow-1;
      end
      ne = ne - 1;
end
% Spherical Cap #2 (Has Hole it top)
if type==5
      nndsx = nelx+1;
   nndsy = nelp+1;
  theta2 = theta2 * pi / 180;
theta1 = theta1 * pi / 180;
alpha2 = alpha2 * pi / 180;
alpha1 = alpha1 * pi / 180;
      tstep = (theta2 - theta1)/nelx;
      astep = (alpha2 - alpha1)/nelp;
      i = 1;
      ne = nelx * nelp;
      nnodes = nndsx * nndsy;
      x = zeros(nnodes, 1);
      y = x;
      z = x;
```

```
nodes = zeros(ne, 4);
      for thetat = thetal:tstep:theta2
            for alphat=alpha1:astep:alpha2
                   x(i) = R * cos(alphat) * cos(thetat);
         y(i) = R * cos(alphat) * sin(thetat);
         z(i) = R * sin(alphat);
                   i = i + 1;
            end
      end
end
% Calculate Element Connectivity (will work for all shapes except cap)
if type~=4
   for i=0:(nelx - 1)
            for j=1:nelp
                   ie = (i * nelp) + j;
nodes(ie,1) = (i * (nelp + 1)) + j;
                   nodes(ie, 2) = nodes(ie, 1) + nelp + 1;
                   nodes(ie,3) = nodes(ie,2) + 1;
                   nodes(ie, 4) = nodes(ie, 1) + 1;
            end
      end
end
% Void content
Phi0 = input('Initial Void Content:');
q1 = input('Value for q1:');
q2 = input('Value for q2 (Enter 0 for Default):');
q3 = input('Value for q3 (Enter 0 for Default):');
fn = input('Nucleating Particle Content:');
en = input('Mean Nucleation Strain (Enter 0 for Default):');
sn = input('Nucleation Standard Deviation (Enter 0 for Default):');
% Create Stress-Strain curve
correct = 'N';
while ((correct == 'n') | (correct == 'N'))
   iseg = input('Number of Plastic Segments (0-9):');
      if (iseg ~= 0)
         for i=1:iseg
            slope(i)=input('Slope of Section:');
            strain(i)=input('Right Strain limit of Section:');
         end
      end
   strn(1) = 0;
   strss(1) = 0;
   strn(2) = Syp/E;
   strss(2) = Syp;
   if (iseg~=0)
      for i=1:iseg
         strn(2+i)=strain(i);
         strss(2+i) = strss(2+i-1) + (strn(2+i) - strn(2+i-1)) * slope(i);
      end
   end
      figure(1);
      plot(strn, strss), grid, xlabel('Strain'), ylabel('Stress'), title('Mat
erial Yield Property')
   correct = input('Is the Stress-Strain Plot Correct? (y or n):','s');
```

```
end
% Fill eprop
eprop=zeros(1,40);
eprop(1) = rho;
eprop(2) = E;
eprop(3) = pois;
eprop(4) = t;
eprop(5) = Syp;
eprop(6) = q1;
eprop(7) = q2;
eprop(8) = q3;
eprop(9) = fn;
eprop(10) = en;
eprop(11) = sn;
eprop(12) = Phi0;
eprop(13) = iseq;
if (iseg > 0)
   for i=1:iseq
      eprop(12+2*i)=slope(i);
      eprop(12+2*i+1)=strain(i);
   end
end
% Display resulting structure
figure(2)
AdjView = [-37.5 \ 30];
shownode;
rotate3d on;
% Apply Boundary Conditions
if type~=4
   bct = zeros(1,nnodes);
   bcr = bct;
end
disp('Boundary Conditions:');
disp('0 = No Constraint');
disp('1 = Constrained in X');
disp('2 = Constrained in Y');
disp('3 = Constrained in Z');
disp('4 = Constrained in X and Y');
disp('5 = Constrained in Y and Z');
disp('6 = Constrained in X and Z');
disp('7 = Constrained in X,Y, and Z');
disp('....');
if type==4,
   disp('(It is inadvisable to change BC''s for Spherical Cap)');
end
nod = input('Node to change Boundary Conditions (0 to end, -1 to
list):');
while ((nod ~= 0) & (nod <= nnodes))
   if nod < 0
      disp('Node
                   Trans
                            Rot');
      for i=1:nnodes
         disp(sprintf('
                                  %d
                                        %d',i,bct(i),bcr(i)));
      end
      disp(sprintf('Current: Translation = %d, Rotation = ',...
```

```
'%d',bct(nod),bcr(nod)));
      bct(nod) = input('New Translation Boundary Condition(0-7):');
      bcr(nod) = input('New Rotational Boundary Condition(0-7):');
      disp(' ');
   nod = input('Node to change Boundary Condition (0 to end):');
end
% Create Load History
disp('(Pressure Load only works for Flat Plate)');
nload = input('Number of nodes to apply load to (0 = pressure load):');
ntime = input('Number of time intervals for Load Description:');
disp('Force Direction:');
disp('1 = Force along X-axis');
disp('2 = Force along Y-axis');
disp('3 = Force along Z-axis');
disp('4 = Moment about X-axis');
disp('5 = Moment about Y-axis');
disp('6 = Moment about Z-axis');
if nload == 0
      nload = nnodes * 6;
      disp('Input Pressure Time History');
      for i=1:ntime
          ldtime(i) = input('Time:');
          Pressure(i) = input('Magnitude(use neg. values for bottom', ...
                          'pressure):');
      end
      F = zeros(nnodes, 1);
      for elem=1:ne
          % Get Coordinates of Nodes for this element
          x1 = x(nodes(elem, 1));
          x2 = x(nodes(elem, 2));
          x3 = x(nodes(elem, 3));
          x4 = x(nodes(elem, 4));
          y1 = y(nodes(elem, 1));
          y2 = y(nodes(elem, 2));
          y3 = y(nodes(elem, 3));
          y4 = y(nodes(elem, 4));
          z1 = z (nodes(elem, 1));
          z2 = z (nodes(elem, 2));
          z3 = z (nodes(elem, 3));
          z4 = z(nodes(elem, 4));
          % Calculate length of sides, then area
          L1 = sqrt((x2-x1)^2+(y2-y1)^2+(z2-z1)^2);
          L2 = sqrt((x3-x2)^2+(y3-y2)^2+(z3-z2)^2);
          L3 = sqrt((x4-x3)^2+(y4-y3)^2+(z4-z3)^2);
          L4 = sqrt((x1-x4)^2+(y1-y4)^2+(z1-z4)^2);
          dot1 = (x1-x2)*(x3-x2)+(y1-y2)*(y3-y2)+(z1-z2)*(z3-z2);
          dot2 = (x1-x4)*(x3-x4)+(y1-y4)*(y3-y4)+(z1-z4)*(z3-z4);
          Area = 0.5 * (sqrt(L1^2 * L2^2 - dot1^2) + sqrt(L3^2 * ...
                    L4^2 - dot2^2);
          % Calculate Force per unit pressure (assumes square elements)
          F(nodes(elem, 1)) = F(nodes(elem, 1)) + Area / 4;
          F(\text{nodes}(\text{elem}, 2)) = F(\text{nodes}(\text{elem}, 2)) + \text{Area} / 4;
          F(\text{nodes}(\text{elem},3)) = F(\text{nodes}(\text{elem},3)) + \text{Area} / 4;
          F(\text{nodes}(\text{elem}, 4)) = F(\text{nodes}(\text{elem}, 4)) + \text{Area} / 4;
      end
```

```
% Create Node, Force, Time Vectors
      nc = 1;
      for n=1:nnodes,
         loadnode(nc) = n;
         loadnode(nc+1) = n;
         loadnode(nc+2) = n;
         loadnode(nc+3) = n;
         loadnode(nc+4) = n;
         loadnode(nc+5) = n;
         loaddir(nc) = 1;
         loaddir(nc+1) = 2;
         loaddir(nc+2) = 3;
         loaddir(nc+3) = 4;
         loaddir(nc+4) = 5;
         loaddir(nc+5) = 6;
         loadtime(nc,:) = ldtime;
         loadtime(nc+1,:) = ldtime;
         loadtime(nc+2,:) = ldtime;
         loadtime(nc+3,:) = ldtime;
         loadtime(nc+4,:) = ldtime;
         loadtime(nc+5,:) = ldtime;
         loadmag(nc,:) = xd(n) * F(n) * Pressure;
         loadmag(nc+1,:) = yd(n) * F(n) * Pressure;
         loadmag(nc+2,:) = zd(n) * F(n) * Pressure;
         loadmag(nc+3,:) = loadmag(nc,:) * t / 2;
         loadmag(nc+4,:) = loadmag(nc+1,:) * t / 2;
         loadmag(nc+5,:) = loadmag(nc+2,:) * t / 2;
         nc = nc + 6;
      end
else
   disp('Input Load/Time/Magnitude History');
   for n=1:nload
      loadnode(n) = input('Node Number:');
      loaddir(n) = input('Load Direction:');
      for nt = 1:ntime
         loadtime(n,nt) = input('Time:');
         loadmag(n,nt) = input('Magnitude:');
      end
   end
end
% Create draft input file
F = fopen('input.in','w');
fprintf(F,'%d 1 %d %d 0 %d 0 \r\n',nnodes,nload,ntime,ipt);
fprintf(F,'%e %e %e %e \r\n', stime, etime, delt, ptime);
fprintf(F,'0 0 0 0 0 0 \r\n');
fprintf(F,'0 %d 0 0 0 \r\n',ne);
fprintf(F,'0 0 0 0 \r\n');
'fprintf(F,'0 0\r\n0 0\r\n%d\r\n1 1 \r\n',ipoint);
for i=1:4
   fprintf(F,'%e %e %e %e %e %e %e %e %e\r\n',...
                   eprop((1+(i-1)*10):(i*10)));
end
for i=1:nnodes
      fprintf(F,'%d %d %d %e %e %e \r\n',...
                   i,bct(i),bcr(i),x(i),y(i),z(i));
end
```

APPENDIX B. POST-PROCESSOR IN MATLAB FOR FEA.

```
% fea disp.m
% P. McDermott
% Uses GUI to read and display output from fea program
Scrn = get(0, 'ScreenSize');
ht = round(0.9 * Scrn(4) / 2 - 50);
wd = round(0.9 * (Scrn(3) - 120) / 2);
hf = figure('NumberTitle','off','Name','FEA','MenuBar','none',...
             'Position', [20 (Scrn(4) - 500) 120 460], 'Color', 'b');
hfs = figure('NumberTitle','off','Name','Structure
View', 'Visib', 'off',...
             'Position', [160 (Scrn(4)-70-ht) wd ht]);
hfd = figure('NumberTitle','off','Name','Displacement','Visib','off',...
             'Position', [(wd+180) (Scrn(4) - 70-ht) wd ht]);
hlast = 0;
AdjView = [-37.5 \ 30];
flagd = 1;
flagv = 0;
flaga = 0;
flags = 0;
File = 'output.fea';
File2 = 'output.fes';
P = pwd;
Path = strcat(P,'\');
hc change = uicontrol(hf,'Style','push','String','Change File',...
             'Position', [20 380 80 20],...
             'CallBack',[...
             '[File1 Path1] = uigetfile(''*.fea'',''Load FEA Output
File'');',..
             'if File1 ~= 0,'...
                   'File = File1;'...
          'Path = Path1;'...
          'File2 = File;'...
          'File2(length(File2)) = ''s'';'...
                   'set(hc file, ''String'', File);'...
             'end']);
hc_load = uicontrol(hf,'Style','check','String','Load File',...
             'Position',[20 420 80 20],...
             'CallBack',[...
          'FilePath = [Path File];',...
          'FilePath2 = [Path File2];',...
                    'parse;',...
                   'set (hc_clear,''Enable'',''on'');',...
'set (hc_disp,''Enable'',''on'');',...
'set (hc_velo,''Enable'',''on'');',...
'set (hc_acc,''Enable'',''on'');',...
                    'set(hc show, ''Enable'', ''on'');',...
```

```
'set(hc_load,''Value'',1,''Enable'',''off'');']);
hc_clear = uicontrol(hf,'Style','push','String','Clear
Data', 'Enable', 'off', ...
             'Position',[20 60 80 20], 'CallBack',[...
             'clear n* x* y* z* i* s* t* v* a* fp factor d etime',...
             'dt m time;',...
'set(hfd,''Visib'',''off'');',...
             'if (flagv>0), set(hfv,''Visib'',''off''); end;',...
             'if (flaga>0), set(hfa,''Visib'',''off''); end;',...
             'set(hfs,''Visib'',''off'');',...
             'flagd = 1; flagv = 0; flaga = 0; flags = 0;',...
             'set(hc load, ''Value'', 0, ''Enable'', ''on'');',...
             'set(hc disp, ''Enable'', ''off'');',...
             'set(hc_velo,''Enable'',''off'');',...
             'set(hc_acc,''Enable'',''off'');',...
             'set(hc show, ''Enable'', ''off'');',...
             'set(hc_replay,''Enable'',''off'');',...
             'hlast = 0;',...
             'set(hc mview, ''Enable'', ''off'');',...
             'set(hc_clear,''Enable'',''off'');']);
hc_close = uicontrol(hf,'Style','push','String','Close',...
                   'Callback',[...
                          'close(hfd);',...
                          'if (flaga>0), close(hfa); end; ',...
                          'if (flagv>0), close(hfv); end;',...
                          'close(hfs);',...
                          'clear;',...
'close;'],...
                   'Position',[20 20 80 20]);
hc_disp = uicontrol(hf,'Style','push','String','Displace','Position',...
             [20 260 80 20],...
                    'Enable','off',...
                    'Callback',[...
                    'if (flags)',...
                          'AdjView = get(hax,''View'');',...
                   'end;',...
                   'figure(hfd);',...
                   'displace;',...
                    'set(hc_replay,''Enable'',''on'');',...
                   'flagd = 0;',...
                   'hlast = hfd;']);
hc mview = uicontrol(hf,'Style','push','String', 'Adjustment',...
                   'Position',[20 100 80 20],...
                    'Enable', 'off',...
                    'CallBack',[...
                          'figure(hfs);',...
                          'mmview3d;']);
hc file = uicontrol(hf, 'Style', 'text', 'Position', [20 340 80 20],...
                   'String', File);
```

```
hc velo = uicontrol(hf,'Style','push','String','Graphs','Position',[20
22\overline{0} 80 20],...
                  'Enable','off',...
          'Callback',[...
            'flagv=1;',...
'control;']);
hc acc = uicontrol(hf, 'Style', 'push', 'String', 'Stress', 'Position', [20]
18\overline{0} 80 20], \dots
                  'Enable', 'off',...
            'Callback',[...
                  'if (flags)',...
'AdjView = get(hax,''View'');',...
            'figure(hfd);',...
'stresscolor;',...
            'flagd = 0;',...
                  'set(hc replay,''Enable'',''on'');',...
                      'hlast = hfd;']);
hc_replay = uicontrol(hf,'Style','push','String','Replay',...
'Position',[20 140 80 20],...
                  'Enable', 'off',...
                  'Callback',[...
                        'figure(hlast);',...
                        'movie(m);']);
'Callback',[...
                        'figure(hfs);',...
                        'shownode;',...
                        'set(hc_mview,''Enable'',''on'');',...
                        'flags = 1;',...
'hax = gca;']);
```

```
% PARSE.M
      This m-file reads the output file generated by fea.f
      and parses it into matlab usable variables which will be used
용
     by another program to graphically display the data in various
      formats
                                  McDermott 1999
% Create Progress Box and 1st Message
hprog = figure('NumberTitle','off','Name','Load File
Progress', 'MenuBar', 'none', ...
   Position', [(Scrn(3)/2 - 80) (Scrn(4)/2 - 40) 260 80], 'Color', 'y');
hmsg = uicontrol('Style', 'text', 'Position',...
      [20 45 220 20], 'BackgroundColor', 'y', ...
   'String', 'Opening Files...');
hmsg2 = uicontrol('Style', 'text', 'Position', ...
      [20 15 220 20], BackgroundColor', 'y', ...
      'String', FilePath);
drawnow;
fp=fopen(FilePath,'r');
fp2 = fopen(FilePath2,'r');
top=fscanf(fp,'%e',33);
% Update Progress Box
set(hmsg,'String','Loading Displacement');
drawnow;
% split into var names
set(hmsq2,'String','Input & eprop');
drawnow;
nnodes=top(1);
nmat=top(2);
nload=top(3);
loadtime=top(4);
initc=top(5);
stime=top(8);
etime=top(9);
dt=top(11);
ne=top(19);
ipoint = top(31);
for i=1:nmat,
      eprop = fscanf(fp,'%e',40);
end;
for i=1:(nnodes),
                                   % dummy index
      fscanf(fp,'%e',3);
                                   % x coordinates of nodes
      x(i)=fscanf(fp,'%e',1);
                                   % y coordinates of nodes
      v(i)=fscanf(fp,'%e',1);
                                   % z coordinates of nodes
      z(i)=fscanf(fp,'%e',1);
end;
```

```
for i=1:ne,
      fscanf(fp,'%e',2);
      nodes(i,:)=fscanf(fp,'%e',4)';
                                            % associate nodes w/ elements
      fscanf(fp,'%e',1);
end;
%initial conditions
fscanf(fp,'%e',((3*(6*nnodes))+(2*nload*(loadtime+1))));
nsteps=round((etime-stime)/dt);
time = [];
temp = nsteps;
for t=1:nsteps,
   set(hmsg2,'String',['Time Step: ' num2str(t) ' / ' num2str(nsteps)]);
   drawnow;
   test=fscanf(fp,'%e',1);
   if (test > 0.0),
      time(t) = test;
             for i=1:nnodes,
             fscanf(fp,'%e',1);
                                            %dummy index
                   xu(i,t)=fscanf(fp,'%e',1);
                   xv(i,t)=fscanf(fp,'%e',1);
                   xa(i,t)=fscanf(fp,'%e',1);
                   fscanf(fp,'%e',1);
                   yu(i,t)=fscanf(fp,'%e',1);
                   yv(i,t)=fscanf(fp,'%e',1);
                   ya(i,t)=fscanf(fp,'%e',1);
                   fscanf(fp,'%e',1);
                   zu(i,t)=fscanf(fp,'%e',1);
                   zv(i,t)=fscanf(fp,'%e',1);
                  za(i,t)=fscanf(fp,'%e',1);
         fscanf(fp,'%e',1);
         thxu(i,t)=fscanf(fp,'%e',1);
         thxv(i,t)=fscanf(fp,'%e',1);
         thxa(i,t)=fscanf(fp,'%e',1);
         fscanf(fp,'%e',1);
         thyu(i,t)=fscanf(fp,'%e',1);
         thyv(i,t)=fscanf(fp,'%e',1);
         thya(i,t)=fscanf(fp,'%e',1);
         fscanf(fp,'%e',1);
         thzu(i,t)=fscanf(fp,'%e',1);
         thzv(i,t)=fscanf(fp,'%e',1);
         thza(i,t)=fscanf(fp,'%e',1);
      end;
   else
      % EOF reached, terminate loop storing maximum number of time steps
      t = nsteps;
   end;
end;
nsteps = size(zu, 2);
fclose(fp);
% Update Progress box
set(hmsg,'String','Calculating Scales');
set(hmsg2,'String',' ');
drawnow;
% Autoscale vectors
dx = abs(max(x) - min(x));
dy = abs(max(y) - min(y));
```

```
dz = abs(max(z) - min(z));
  % Check for flat structures
  if dx == 0
         dx = 1;
  end
  if dy == 0
         dy = 1;
  end
  if dz == 0
         dz = 1;
  end
  scale(1) = dx / max(max(abs(xu)));
scale(2) = dy / max(max(abs(yu)));
  scale(3) = dz / max(max(abs(zu)));
  dscale = min(scale) * 0.25;
  if dscale < 1
         dscale = 1;
  end
  xu = xu * dscale;
  yu = yu * dscale;
  zu = zu * dscale;
  scale(1) = dx / max(max(abs(xv)));
  scale(2) = dy / max(max(abs(yv)));
  scale(3) = dz / max(max(abs(zv)));
  vscale = min(scale) * 0.75;
  if vscale < 1
         vscale = 1;
  end
xv = xv * vscale;
  yv = yv * vscale;
  zv = zv * vscale;
  scale(1) = dx / max(max(abs(xa)));
scale(2) = dy / max(max(abs(ya)));
scale(3) = dz / max(max(abs(za)));
ascale = min(scale) * 0.5;
  if ascale < 1
         ascale = 1;
  end
  xa = xa * ascale;
 'ya = ya * ascale;
  za = za * ascale;
```

```
% Allocate arrays for stress-strain data
stress = zeros(ipoint*ne*6,nsteps);
strain = stress;
plastrn = stress;
void = zeros(ipoint*ne,nsteps);
yieldstrs = void;
effplstrn = void;
vmstress = void;
effstrain = void;
estrain = void;
%pstrain = void;
% Update Progress Box
set(hmsg, 'String', 'Loading Stress & Strain');
drawnow;
% Read Stress-Strain Data
for t=1:nsteps,
   test = fscanf(fp2,'%s',1);
   set(hmsq2,'String',['Time Step: ' num2str(t) ' / ' num2str(nsteps)]);
   drawnow;
   if (test > 0.0),
      t1=fscanf(fp2, '%e',1);
      for elem=1:ne,
         fscanf(fp2,'%s',1);
             fscanf(fp2,'%d',1);
         for point=1:ipoint
             index = (elem-1)*6*ipoint + (point-1)*6;
             fscanf(fp2,'%d',1);
             for i = 1:6,
                ind = i + index;
                stress(i + index,t)=fscanf(fp2,'%e',1);
                strain(i + index,t)=fscanf(fp2,'%e',1);
                plastrn(i + index,t)=fscanf(fp2,'%e',1);
             end
             index1 = (elem-1)*ipoint + point;
             yieldstrs(index1,t) = fscanf(fp2,'%e',1);
             void(index1,t) = fscanf(fp2,'%e',1);
             effplstrn(index1,t) = fscanf(fp2,'%e',1);
         end
      end
   else
      t = nsteps;
   end
end
% Calculate Von-Misses Stress and an equivalent total strain value
% Update Progress Box
set(hmsg,'String','Calculating Effective');
set(hmsg2,'String','Stress and Strain...');
drawnow;
fclose(fp2);
% Adjust for total strain
%totalstrn = strain + plastrn;
```

```
% Get Effective stress and strain (doesn't work with multiple materials)
for elem=1:ne,
  for point=1:ipoint,
     index = (elem-1)*6*ipoint + (point-1)*6 + 1;
     index1 = (elem-1)*ipoint + point;
     vmstress(index1,:)=sqrt(0.5*((stress(index,:) - ...
           stress(index+1,:)).^2 + ...
           ( stress(index+1,:) - stress(index+2,:)).^2 +...
           ( stress(index+2,:) - stress(index,:)).^2 + 6 *...
           (stress(index+3,:).^2 + stress(index+4,:).^2 ...
           + stress(index+5,:).^2)));
     estrain(index1,:)=sqrt(2*((strain(index,:)-strain(index+1,:)).^2...
           +(strain(index+1,:)-strain(index+2)).^2 + (strain(index+2)...
           - strain(index,:)).^2) + 3*(strain(index+3).^2 + ...
           strain(index+4).^2 + strain(index+5).^2))/(2*(1+eprop(3)));
     end
end
close (hprog);
```

```
% Control.m
      Display Von-Misses Stress and Yield Stress curves
                                                  McDermott 1999
hfv = figure('NumberTitle','off','Name','Graph Control','Visib','on',...
             'MenuBar', 'none', 'Color', 'b', 'Position', [160 50 wd ht]);
hfa = figure('NumberTitle','off','Name','Plot Window','Visib','off',...
             'Position', [(wd+180) 50 wd ht]);
elementstr = ['1|'];
for i=2:ne
   elementstr = strcat(elementstr, num2str(i));
   elementstr = strcat(elementstr,'|');
end
pointstr = ['1|'];
for i=2:ipoint
   pointstr = strcat(pointstr,num2str(i));
   pointstr = strcat(pointstr,'|');
end
nodestr = ['1|'];
for i=2:nnodes
   nodestr = strcat(nodestr,num2str(i));
   nodestr = strcat(nodestr,'|');
end
typestr = ['Stress vs Strain|Stress vs Time|Strain vs time|Porosity vs
Time|',...
      'Porosity vs Strain|Sigma zz vs Thickness|Sigma xx vs Time|',...
      'Sigma yy vs Time|Sigma zz vs Time|Sigma Multiplot|',...
      'X Disp. vs Time|Y Disp. vs Time|Z Disp. vs Time|',...
      'Theta x vs Time|Theta y vs Time|',...
      'Theta z vs Time | MeanDisp vs Time | Plastic Strain | Elastic Strain | ];
% Start at Bottom
hv_plot = uicontrol(hfv,'Style','push','Position',...
      [(wd/2 - 40) 20 80 20], 'String', 'Plot', 'Callback', ...
      'flaga=1;drawstress;');
hv_typetxt = uicontrol(hfv,'Style','text','Position',[ 20 60 120 20],...
   'String', 'Graph Type:');
hv type = uicontrol(hfv, 'Style', 'popupmenu', 'Position', ...
            [ 160 60 160 20 ], 'String', typestr, 'Value', 1, ...
             'Callback', 'type = get(hv_type, ''Value'');');
type = 1;
hv pointtxt = uicontrol(hfv,'Style','text','Position',...
             [ 20 100 120 20 ], 'String', 'Integration Point:');
hv_point = uicontrol(hfv,'Style','popupmenu','Position',...
[ 280 100 40 20 ],'String',pointstr,'Value',1,...
             'Callback', 'point = get(hv_point, ''Value'');');
point = 1;
hv elemtxt = uicontrol(hfv, 'Style', 'text', 'Position', ...
            [ 20 140 120 20 ], 'String', 'Element Number:');
```

```
hv elem = uicontrol(hfv,'Style','popupmenu','Position',...
            [280 140 40 20 ], 'String', elementstr, 'Value', 1, ...
            'Callback', 'selem = get(hv_elem, ''Value'');');
selem = 1;
hv nodetxt = uicontrol(hfv,'Style','text','Position',...
            [20 180 120 20 ], 'String', 'Node Number:');
hv_node = uicontrol(hfv,'Style','popupmenu','Position',...
            [280 180 40 20], 'String', nodestr, 'Value', 1, ...
            'Callback', 'snode = get(hv_node, ''Value'');');
snode = 1;
hv slidtxt = uicontrol(hfv,'Style','text','Position',[20 260 120 20],...
   'String', 'Time Slider Value: ');
hv slidtxt2 = uicontrol(hfv,'Style','text','Position',...
            [160 260 80 20], 'String', '0');
hv_slidtxt3 = uicontrol(hfv,'Style','text','Position',...
            [240 260 80 20], 'String', ' Seconds');
hv slid = uicontrol(hfv,'Style','slider','Position',...
            [20 220 (wd - 40) 20], 'Min', 1, 'Max', nsteps, 'Value', 1, ...
            'Callback',...
      ['tstp = round(get(hv_slid,''Value''));',...
       'set(hv slidtxt2, ''String'', num2str(time(tstp)));']);
tstp = 1;
set(hfv,'Visib','on');
```

```
% displace.m
      Display the element and its displacement at each time step
                                              McDermott 1999
% plot the element(s)
if flagd
      zmax=max(z)+max(max(zu))+.1;
      ymax=max(y)+max(max(yu))+.1;
      xmax=max(x)+max(max(xu))+.1;
      zmin=min(z)+min(min(zu))-.1;
      ymin=min(y)+min(min(yu))-.1;
      xmin=min(x)+min(min(xu))-.1;
      AxMax = [xmin, xmax, ymin, ymax, zmin, zmax];
end;
m=moviein(nsteps);
% Display scaling factor in title bar
dispstring = ['Displacement: Scale = ' num2str(dscale,'%1.4g')];
set(gcf,'Name',dispstring);
for t=1:nsteps
      % display global node numbers
      cla,xlabel('x axis'),ylabel('y axis'),zlabel('Z axis'),grid on,...
      set(gca,'Box','off'),hold on;
      for i=1:ne
           x1=x (nodes(i,1))+xu (nodes(i,1),t);
           x2=x (nodes(i,2))+xu (nodes(i,2),t);
           x3=x (nodes(i,3))+xu (nodes(i,3),t);
           x4=x (nodes(i,4))+xu (nodes(i,4),t);
           y1=y(nodes(i,1))+yu(nodes(i,1),t);
           y2=y (nodes(i,2))+yu (nodes(i,2),t);
           y3=y (nodes (i,3))+yu (nodes (i,3),t);
           y4=y (nodes(i,4))+yu (nodes(i,4),t);
           z1=z (nodes(i,1))+zu(nodes(i,1),t);
           z2=z (nodes(i,2))+zu (nodes(i,2),t);
           z3=z (nodes(i,3))+zu(nodes(i,3),t);
           z4=z (nodes(i,4))+zu(nodes(i,4),t);
           xvec=[x1, x2, x3, x4, x1];
           yvec=[y1, y2, y3, y4, y1];
           zvec=[z1, z2, z3, z4, z1];
           plot3(xvec, yvec, zvec), view(AdjView), axis(AxMax);
      end;
      hold off;
     m(:,t)=getframe;
end;
movie(m);
```

```
% shownode.m
     Display the element
용
                                 McDermott 1999
% Remove previous contents
cla;
% plot the element(s)
offset = 0.1 * (max(z) + max(y) + max(x))/3;
zmax=max(z)+offset;
ymax=max(y)+offset;
xmax=max(x)+offset;
zmin=min(z)-offset;
ymin=min(y)-offset;
xmin=min(x)-offset;
AxMax = [xmin, xmax, ymin, ymax, zmin, zmax];
xlabel('x axis'), ylabel('y axis'), zlabel('z axis'),...
   axis (AxMax), view (AdjView), grid, hold on;
for i=1:ne
     x1=x(nodes(i,1));
     x2=x(nodes(i,2));
     x3=x(nodes(i,3));
     x4=x(nodes(i,4));
     y1=y(nodes(i,1));
     y2=y(nodes(i,2));
     y3=y(nodes(i,3));
     y4=y(nodes(i,4));
     z1=z (nodes(i,1));
     z2=z (nodes(i,2));
     z3=z (nodes(i,3));
     z4=z (nodes(i,4));
     xvec=[x1, x2, x3, x4, x1];
     yvec=[y1, y2, y3, y4, y1];
     zvec=[z1, z2, z3, z4, z1];
     plot3(xvec, yvec, zvec);
end;
d=offset * 0.25;
for i=1:nnodes
     text(x(i)+d, y(i)+(2*d),z(i)+d,num2str(i));
end
```

```
% drawstress.m - plots Von-Mises stress and Yield stress
                                          McDermott 1999
figure(hfa);
set(hfa,'Visib','on');
flaga = 1;
% tstp = time vector index (not all plots need)
% point = selected integration point
% selem = selected element
% type = Graph Type selected:
            1 = Stress vs Strain
            2 = Stress vs Time
용
            3 = Strain vs Time
ક્ષ
            4 = Porosity vs Time
            5 = Porosity vs Plastic Strain
            6 = Z Stress vs Thickness (at time(tstp))
용
            7 = X Stress vs Time
            8 = Y Stress vs Time
            9 = Z Stress vs Time
용
            10 = Stress Multiplot (all 4 stress plots vs Time)
용
용
            11 = X Displacement vs Time
용
            12 = Y Displacement vs Time
            13 = Z Displacement vs Time
용
            14 = Theta X vs Time
용
            15 = Theta Y vs Time
용
            16 = Theta Z vs Time
용
            17 = Mean Displacement vs Time
용
            18 = Plastic Strain vs Time
용
            19 = Elastic Strain vs Time
index = (selem-1)*6*ipoint + (point-1)*6;
index1 = (selem-1)*ipoint + point;
switch type
      case 1,
            titlestr = ['Element #' num2str(selem),...
                  ' Integration Point #' num2str(point)];
plot(effstrain(index1,:), vmstress(index1,:), effstrain(index1,:),...
         yieldstrs(index1,:),'--'),grid,xlabel('Effective Strain'),...
         ylabel('Effective Stress'), title(titlestr), ...
         legend('VME Stress','Yield Stress')
      maxvm = max(vmstress(index1,:))
      maxstrn = max(effstrain(index1,:))
   case 2,
      titlestr = ['Element #' num2str(selem) ' Integration Point #',...
             num2str(point)];
      plot(time, vmstress(index1,:), time, yieldstrs(index1,:),...
            '--'), grid, xlabel('Time (sec)'), ...
            ylabel('Effective Stress'),title(titlestr),...
            legend('VME Stress', 'Yield Stress');
      maxvm = max(vmstress(index1,:))
      maxys = max(yieldstrs(index1,:))
```

```
case 3,
    titlestr = ['Element #' num2str(selem) ' Integration Point #',...
                num2str(point)];
    plot(time,effstrain(index1,:)),grid,xlabel('Time (sec)'),...
       ylabel('Effective Strain'), title(titlestr);
    maxstrn = max(effstrain(index1,:))
case 4,
    titlestr = ['Element #' num2str(selem) ' Integration Point #',...
                num2str(point)];
    plot(time, void(index1,:)), grid, xlabel('Time (sec)'), ...
                title(titlestr), ylabel('Porosity');
   maxpor = max(void(index1,:))
 case 5,
    titlestr = ['Element #' num2str(selem) ' Integration Point #',...
                num2str(point)];
    plot(effplstrn(index1,:),void(index1,:)),grid,...
          xlabel('Effective Plastic Strain'),...
          vlabel('Porosity'), title(titlestr);
    maxpor = max(void(index1,:))
 case 6.
    titlestr = ['Element #' num2str(selem) ' Time: ',...
          num2str(time(tstp)) ' sec'];
    strs = zeros(ipoint,1);
    pnts = 1:ipoint;
    for p=1:ipoint
          index3 = (selem-1)*6*ipoint + (p-1)*6 + 3;
          strs(p) = stress(index3,tstp);
    end
    maxstrs = max(strs(p))
    minstrs = min(strs(p))
    plot(strs,pnts),grid,axis ij,ylabel('Integration Point'),...
          xlabel('Z Stress'), title(titlestr);
 case 7,
    titlestr = ['Element #' num2str(selem) ' Integration Point #',...
          num2str(point)];
    plot(time, stress(index+1,:)), grid, xlabel('Time (sec)'), ...
          ylabel('X Stress'), title(titlestr);
    maxstrs = max(stress(index+1,:))
    minstrs = min(stress(index+1,:))
    titlestr = ['Element #' num2str(selem) ' Integration Point #',...
          num2str(point)];
    plot(time, stress(index+2,:)), grid, xlabel('Time (sec)'), ...
          ylabel('Y Stress'), title(titlestr);
    maxstrs = max(stress(index+2,:))
    minstrs = min(stress(index+2,:))
    titlestr = ['Element #' num2str(selem) ' Integration Point #',...
          num2str(point)];
    plot(time, stress(index+3,:)), grid, xlabel('Time (sec)'), ....
          ylabel('Z Stress'), title(titlestr);
    maxstrs = max(stress(index+3,:))
    minstrs = min(stress(index+3,:))
```

```
case 10,
   titlestr = ['Element #' num2str(selem) ' Integration Point #',...
         num2str(point)];
   subplot(2,2,1)
   plot(time, stress(index+1,:)), grid, xlabel('Time (sec)'), ...
         vlabel('X Stress'), title(titlestr);
   subplot(2,2,2)
   plot(time, stress(index+2,:)), grid, xlabel('Time (sec)'),...
         vlabel('Y Stress'), title(titlestr);
   subplot(2,2,3)
   plot(time, stress(index+3,:)), grid, xlabel('Time (sec)'), ...
         ylabel('Z Stress'), title(titlestr);
   subplot(2,2,4)
   plot(time, vmstress(index1,:), time, yieldstrs(index1,:), ...
         '--'), grid, xlabel('Time (sec)'), ...
         ylabel('Effective Stress'), ...
         legend('VME Stress', 'Yield Stress'), title(titlestr);
  maxstrs x = max(stress(index+1,:))
  minstrs x = min(stress(index+1,:))
  maxstrs y = max(stress(index+2,:))
  minstrs_y = min(stress(index+2,:))
  maxstrs z = max(stress(index+3,:))
  minstrs z = min(stress(index+3,:))
case 11,
   titlestr = ['Node #',num2str(snode)];
  plot(time, xu(snode,:)/dscale), grid, xlabel('Time (sec)'),...
         ylabel('X Node Displacement'), title(titlestr);
  maxdx = max(xu(snode,:))/dscale
  mindx = min(xu(snode,:))/dscale
case 12,
  titlestr = ['Node #',num2str(snode)];
  plot(time, yu(snode,:)/dscale), grid, xlabel('Time (sec)'), ...
         ylabel('Y Node Displacement'), title(titlestr);
  maxdy = max(yu(snode,:))/dscale
  mindy = min(yu(snode,:))/dscale
case 13.
  titlestr = ['Node #', num2str(snode)];
  plot(time, zu(snode,:)/dscale), grid, xlabel('Time (sec)'), ...
         ylabel('Z Node Displacement'), title(titlestr);
  maxdz = max(zu(snode,:))/dscale
  mindz = min(zu(snode,:))/dscale
case 14,
  titlestr = ['Node #',num2str(snode)];
  plot(time, thxu(snode,:)), grid, xlabel('Time (sec)'), ...
         ylabel('Theta X'), title(titlestr);
  maxdz = max(thxu(snode,:))
  mindz = min(thxu(snode,:))
   titlestr = ['Node #', num2str(snode)];
  plot(time,thyu(snode,:)),grid,xlabel('Time (sec)'),...
         ylabel('Theta Y'), title(titlestr);
  maxdz = max(thyu(snode,:))
  mindz = min(thyu(snode,:))
```

```
case 16,
     titlestr = ['Node #', num2str(snode)];
     plot(time,thzu(snode,:)),grid,xlabel('Time (sec)'),...
           ylabel('Theta Z'), title(titlestr);
     maxdz = max(thzu(snode,:))
     mindz = min(thzu(snode,:))
  case 17,
     titlestr = ['Node #', num2str(snode)];
     du = sqrt(xu(snode,:).^2 + yu(snode,:).^2 + ...
           zu(snode,:).^2)/dscale;
     maxdu = max(du)
     mindu = min(du)
     plot(time,du),grid,xlabel('Time (sec)'),...
           ylabel('Mean Node Displacement'), title(titlestr);
  case 18,
     titlestr = ['Element #' num2str(selem) ' Integration Point #',...
           num2str(point)];
     plot(time, effplstrn(index1,:)), grid, xlabel('Time (sec)'), ...
              ylabel('Effective Plastic Strain'), title(titlestr);
     maxplaststrn = max(effplstrn(index1,:))
  case 19,
     titlestr = ['Element #', num2str(selem) ' Integration Point #',...
           num2str(point)];
     plot(time, estrain(index1,:)), grid, xlabel('Time (sec)'),...
        ylabel('Effective Elastic Strain'), title(titlestr);
     maxelaststrn = max(estrain(index1,:))
  otherwise,
     disp('Invalid Graph Type');
end
```

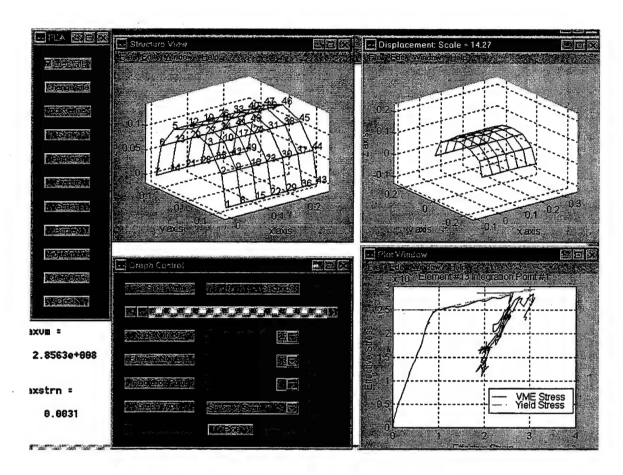


Figure 23. Screen Capture of FEA Preprocessor.

APPENDIX C. FILE LISTING FOR FEA SHELL ELEMENT SUBROUTINE.

```
c*
      elsh41.f
c*
      Contains the following subroutines:
c*
            elsh4l - element matrices for 3-D degenerated 4-node shell
          *****************
      subroutine elsh41 (nnds, xcoord, ycoord, zcoord, nesh4, ndsh4, matsh4,
     1
                      nmat, matno, eprop, ndsdof, sdisp, svel, sforce,
     2
                       smass, nsdof, ishel4, stssh4, mpoint, ipoint, itime)
  ********************
      compute and assemble element matrices for shell elements with
C
      4 nodes (linear analysis)
C
      3-D degenerated formulation including transverse shear and
С
      transverse normal deformation
С
      pressure formulation
                                             McDermott 1999
C**********************
      common /fblk/ fail, array1(125)
      dimension xcoord(nnds), ycoord(nnds), zcoord(nnds),
                ndsh4(nesh4+1,4), matsh4(nesh4+1), ishel4(nesh4+1),
     2
                matno(nmat), eprop(nmat, 40), ndsdof(nnds),
     3
                sdisp(nsdof), svel(nsdof), sforce(nsdof), smass(nsdof),
                stssh4 (mpoint, nesh4*8)
      dimension eforce (24), lnode (4), evel (24), slope (10), dstrnplst (6),
                emass(24), gausx(4), lindex(24), strain(10), corr(2),
     1
     2
                array2(15), estrain(6), estress(6), s(6), A(2,2), Ainv(2,2),
     3
                gausy(4), weighx(4), weighy(4), shape1(3), deriv1(3),
     4
                shapef(4), deriv1(2,4), derivg(3,4), aj(3,3), ajinv(3,3),
     5
                bmtx(6,24), tmp(8), xc(4), yc(4), zc(4), estressg(6),
     6
                derilg(3,4),d1(3),edisp(24),edispg(24),rot6t(6,6),
     7
                d2(3), v1(3), v2(3), v3(3), gausz(10), weighz(10),
     8
                bmtxt(24,6),delt(6),t1(3,3),tlinv(3,3),B(2),
     9
                rot6(6,6),rot3(3,3),estrainp(6),eforceg(24),
     1
                weighxh(4), weighyh(4), weighzh(4), eforceh(24),
                gausxh(4), gausyh(4), gauszh(4), iplane(5)
С
      data ngausx, ngausy, ndof, n /1, 1, 6, 4/
      data ngausxh, ngausyh, ngauszh, hour1 /2, 2, 1, 0.05/
      data ref,eps /0.0,1.0e-4 /
      data delt /1.0, 1.0, 1.0, 0.0, 0.0, 0.0/
      data iplane /1, 2, 4, 5, 6/
     neldof=ndof*n
      ngausz=ipoint
      ngauss=ngausx*ngausy*ngausz
      ngaussh=ngausxh*ngausyh*ngauszh
  *** extract gauss points and weights for numerical integration
С
      call gauss (ngausx, gausx, weighx)
      call gauss (ngausy, gausy, weighy)
      call gauss (ngausz, gausz, weighz)
      call gauss (ngausxh, gausxh, weighxh)
      call gauss (ngausyh, gausyh, weighyh)
      call gauss (ngauszh, gauszh, weighzh)
```

```
*** loop for number of elements
С
       do 370 ie=1, nesh4
С
             igausp=0
C
C
  *** initialize element internal force vector
             do 10 i=1, neldof
              eforceh(i) = 0.0
               eforce(i) = 0.0
  10
             continue
C
  *** extract material property index and material properties
С
             imt=matsh4(ie)
             mattyp=matno(imt)
             densty=eprop(imt,1)
             elast = eprop(imt,2)
             poiss = eprop(imt,3)
             thick = eprop(imt, 4)
       Syp = eprop(imt, 5)
       q1 = eprop(imt, 6)
       q2 = eprop(imt, 7)
      q3 = eprop(imt, 8)
      fn = eprop(imt, 9)
      en = eprop(imt, 10)
       sn = eprop(imt, 11)
       Phi0 = eprop(imt, 12)
      iseg = eprop(imt, 13) + 0.1
       ystrain = Syp / elast
       strain(1) = ystrain
             if (iseg.gt.0) then
             do 20 i=1,iseg
                    slope(i) = eprop(imt, 12 + i * 2)
                    strain(i+1) = eprop(imt, 13 + i * 2)
   20
             continue
       endif
             bulk = elast / (3.0 * (1.0 - 2.0 * poiss))
shear = elast / (2.0 * (1.0 + poiss))
  *** Default Values
       if (ql .ne. 0.0) then
             if (q2 .eq. 0.0) q2 = 1.0
if (q3 .eq. 0.0) q3 = q1**2
if (fn .ne. 0.0) then
                    if (sn .eq. 0.0) sn = 0.1
                    if (en .eq. 0.0) en = 0.3
             endif
       endif
  *** find dof index of each element
       do 30 i=1,n
             lnode(i)=ndsh4(ie,i)
             do 30 j=1, ndof
                    lindex((i-1)*ndof+j)=ndsdof(lnode(i))+j-1
   30
             continue
```

```
*** load displacement for element
             do 40 i=1, neldof
             edispg(i) = sdisp(lindex(i))
             evel(i) = svel(lindex(i))
   40
             continue
С
C
  *** extract element nodal coordinate values
С
            do 50 i=1,n
            xc(i)=xcoord(lnode(i))
            yc(i)=ycoord(lnode(i))
            zc(i)=zcoord(lnode(i))
   50
            continue
C
c *** compute direction vectors
c *** (v1 & v2: tangent, v3: normal to surface)
            d1(1) = xc(2) - xc(1)
            d1(2) = yc(2) - yc(1)
            d1(3) = zc(2) - zc(1)
            d2(1) = xc(4) - xc(1)
            d2(2) = yc(4) - yc(1)
            d2(3) = zc(4) - zc(1)
  *** Calculation of Hourglass Control Multiplier
С
С
      Need Surface Area of Element
C
      all=sqrt(d1(1)**2+d1(2)**2+d1(3)**2)
      al4 = sqrt(d2(1)**2+d2(2)**2+d2(3)**2)
      al2 = sqrt((xc(3)-xc(2))**2+(yc(3)-yc(2))**2+(zc(3)-zc(2))**2)
      al3 = sqrt((xc(3)-xc(4))**2+(yc(3)-yc(4))**2+(zc(3)-zc(4))**2)
      adot1 = (xc(1)-xc(2))*(xc(3)-xc(2))+(yc(1)-yc(2))*(yc(3)-yc(2))
    1 + (zc(1)-zc(2))*(zc(3)-zc(2))
      adot2 = (xc(1)-xc(4))*(xc(3)-xc(4))+(yc(1)-yc(4))*(yc(3)-yc(4))
    1 + (zc(1)-zc(4))*(zc(3)-zc(4))
      area = 0.5*(sqrt(al1**2*al2**2-adot1**2)+
    1 sqrt(al3**2*al4**2-adot2**2))
      hour = hour1 * thick**2 / area
c *** v3 is normal to plane
            v3(1)=d1(2)*d2(3)-d1(3)*d2(2)
            v3(2)=d1(3)*d2(1)-d1(1)*d2(3)
            v3(3)=d1(1)*d2(2)-d1(2)*d2(1)
            sum = sqrt(v3(1)**2+v3(2)**2+v3(3)**2)
            v3(1) = v3(1) / sum
            v3(2)=v3(2)/sum
            v3(3)=v3(3)/sum
  *** v2 is unit vector in direction of node 1 to node 4
            v2(1) = d2(1)/a14
            v2(2) = d2(2)/a14
            v2(3) = d2(3)/a14
c *** v1 is orthogonal to v2, in plane normal to v3
            v1(1)=v3(2)*v2(3)-v3(3)*v2(2)
            v1(2) = v3(3) * v2(1) - v3(1) * v2(3)
            v1(3)=v3(1)*v2(2)-v3(2)*v2(1)
            sum = sqrt(v1(1)**2+v1(2)**2+v1(3)**2)
```

```
v1(1) = -v1(1) / sum
             v1(2) = -v1(2) / sum
             v1(3) = -v1(3) / sum
 c *** Strain Transformation Matrix
             rot6(1,1)=v1(1)**2
             rot6(1,2)=v1(2)**2
             rot6(1,3)=v1(3)**2
             rot6(1,4)=v1(1)*v1(2)
             rot6(1,5)=v1(2)*v1(3)
             rot6(1,6)=v1(1)*v1(3)
             rot6(2,1)=v2(1)**2
             rot6(2,2)=v2(2)**2
             rot6(2,3)=v2(3)**2
             rot6(2,4)=v2(1)*v2(2)
             rot6(2,5)=v2(2)*v2(3)
             rot6(2,6)=v2(1)*v2(3)
             rot6(3,1)=v3(1)**2
             rot6(3,2)=v3(2)**2
             rot6(3,3)=v3(3)**2
             rot6(3,4)=v3(1)*v3(2)
             rot6(3,5)=v3(2)*v3(3)
             rot6(3,6)=v3(1)*v3(3)
             rot6(4,1)=2.0*v1(1)*v2(1)
             rot6(4,2)=2.0*v1(2)*v2(2)
             rot6(4,3)=2.0*v1(3)*v2(3)
             rot6(4,4)=v1(1)*v2(2)+v2(1)*v1(2)
             rot6(4,5)=v1(2)*v2(3)+v2(2)*v1(3)
             rot6(4,6)=v1(3)*v2(1)+v2(3)*v1(1)
             rot6(5,1)=2.0*v2(1)*v3(1)
             rot6(5,2)=2.0*v2(2)*v3(2)
             rot6(5,3)=2.0*v2(3)*v3(3)
             rot6(5,4)=v2(1)*v3(2)+v3(1)*v2(2)
             rot6(5,5)=v2(2)*v3(3)+v3(2)*v2(3)
             rot6(5,6)=v2(3)*v3(1)+v3(3)*v2(1)
             rot6(6,1)=2.0*v3(1)*v1(1)
             rot6(6,2)=2.0*v3(2)*v1(2)
             rot6(6,3)=2.0*v3(3)*v1(3)
             rot6(6,4)=v3(1)*v1(2)+v1(1)*v3(2)
             rot6(6,5)=v3(2)*v1(3)+v1(2)*v3(3)
             rot6(6,6)=v3(3)*v1(1)+v1(3)*v3(1)
C
   *** Transpose of Transformation Matrix
· C
С
             do 60 i=1,6
             do 60 j=1,6
                   rot6t(i,j)=rot6(j,i)
    60
             continue
 C
   *** Rotation tranformation matrix, and its inverse
 С
             do 70 i=1,3
             t1(i,1) = v1(i)
             t1(i,2) = v2(i)
             t1(i,3) = v3(i)
    70
             continue
             call inv3x3(t1,t1inv,det)
   *** Transform Rotation displacement to local coordinates
```

```
do 80 i=1,4
                                            do 80 j=1,3
                                                                   edisp((i-1)*6+j)=edispg((i-1)*6+j)
                                                                   edisp((i-1)*6+3+j)=edispg((i-1)*6+4)*t1inv(j,1)
                                                                                                               +edispg((i-1)*6+5)*tlinv(j,2)
                  2
                                                                                                                                 +edispg((i-1)*6+6)*tlinv(j,3)
           80
                                             continue
       *** Begining of Hourglass control
                      if (hour.gt.0.0) then
                          do 140 ix=1,ngausxh
                             rc=gausxh(ix)
                             wx=weighxh(ix)
                                 do 140 iy=1, ngausyh
                                     sc=gausyh(iy)
                                     wy=weighyh(iy)
                                        do 140 iz=1, ngauszh
                                            tc=gauszh(iz)
                                            wz=weighzh(iz)
       *** compute shape functions and derivatives
С
                                            call shp2d4 (rc,sc,shapef)
                                            call der2d4 (rc,sc,derivl)
                                            call shp1d2 (tc, shape1)
                                            call der1d2 (tc,deriv1)
C
С
       *** compute jacobian matrix and its inverse
                                            hz=thick*0.5*(shape1(2)*(1.0-ref)-shape1(1)*(1.0+ref))
                                            hzdt=thick*0.5
                     aj(1,1) = derivl(1,1) *xc(1) + derivl(1,2) *xc(2) + derivl(1,3) *xc(3) +
                                                                                        derivl(1,4)*xc(4)+derivl(1,1)*hz*v3(1)+
                  1
                  2
                                                                                        derivl(1,2)*hz*v3(1)+derivl(1,3)*hz*v3(1)+
                  3
                                                                                        derivl(1,4)*hz*v3(1)
                     aj(2,1) = derivl(2,1) *xc(1) + derivl(2,2) *xc(2) + derivl(2,3) *xc(3) + derivl(3,3) *xc(3,3) *xc(3) + derivl(3,3) *xc(3) + derivl(3,
                                                                                        derivl(2,4)*xc(4)+derivl(2,1)*hz*v3(1)+
                  2
                                                                                        derivl(2,2)*hz*v3(1)+derivl(2,3)*hz*v3(1)+
                  3
                                                                                        derivl(2,4)*hz*v3(1)
                                            ai(3,1) = hzdt*v3(1)
                     aj(1,2) = derivl(1,1)*yc(1)+derivl(1,2)*yc(2)+derivl(1,3)*yc(3)+
                  1
                                                                                        derivl(1,4)*yc(4)+derivl(1,1)*hz*v3(2)+
                  2
                                                                                        derivl(1,2)*hz*v3(2)+derivl(1,3)*hz*v3(2)+
                                                                                        derivl(1,4)*hz*v3(2)
                  3
                     aj(2,2) = derivl(2,1) *yc(1) + derivl(2,2) *yc(2) + derivl(2,3) *yc(3) +
                                                                                        derivl(2,4)*yc(4)+derivl(2,1)*hz*v3(2)+
                  2
                                                                                        deriv1(2,2)*hz*v3(2)+deriv1(2,3)*hz*v3(2)+
                  3
                                                                                        derivl(2,4)*hz*v3(2)
                                            aj(3,2) = hzdt*v3(2)
                     aj(1,3) = derivl(1,1)*zc(1)+derivl(1,2)*zc(2)+derivl(1,3)*zc(3)+derivl(1,3)*zc(3)+derivl(1,3)*zc(3)+derivl(1,3)*zc(3)+derivl(1,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+derivl(3,3)*zc(3)+
                                                                                        derivl(1,4)*zc(4)+derivl(1,1)*hz*v3(3)+
                  1
                  2
                                                                                        derivl(1,2)*hz*v3(3)+derivl(1,3)*hz*v3(3)+
                  3
                                                                                        derivl(1,4)*hz*v3(3)
```

```
ai(2,3) = deriv1(2,1)*zc(1)+deriv1(2,2)*zc(2)+deriv1(2,3)*zc(3)+
                          derivl(2,4)*zc(4)+derivl(2,1)*hz*v3(3)+
     2
                          deriv1(2,2)*hz*v3(3)+deriv1(2,3)*hz*v3(3)+
     3
                          derivl(2,4)*hz*v3(3)
             aj(3,3) = hzdt*v3(3)
С
             call inv3x3 (aj,ajinv,det)
  *** compute global derivatives and strain-nodal displacement matrix
С
             do 90 i=1,n
      derivg(1,i)=ajinv(1,1)*derivl(1,i)+ajinv(1,2)*derivl(2,i)
      derivg(2,i)=ajiny(2,1)*derivl(1,i)+ajinv(2,2)*derivl(2,i)
      derivg(3,i)=ajinv(3,1)*derivl(1,i)+ajinv(3,2)*derivl(2,i)
                   derilg(1,i)=ajinv(1,3)*hzdt
                   derilg(2,i)=ajinv(2,3)*hzdt
                   derilg(3,i)=ajinv(3,3)*hzdt
   90
             continue
             do 100 i=1,6
                   do 100 j=1,24
                          bmtx(i,j)=0.0
  100
             continue
             do 110 i=1,n
                   i1=(i-1)*6+1
                   i2=i1+1
                   i3=i2+1
                   i4=i3+1
                   i5 = i4 + 1
                   i6=i5+1
                   gkl=derivg(1,i)*hz+shapef(i)*derilg(1,i)
                   gk2=derivg(2,i)*hz+shapef(i)*derilg(2,i)
                   gk3=derivg(3,i)*hz+shapef(i)*derilg(3,i)
C
                   bmtx(1,i1) = derivg(1,i)
                   bmtx(1,i4)=gk1*(-v2(1))
                   bmtx(1, i5) = gk1*v1(1)
                   bmtx(1, i6) = gk1*v3(1)
                   bmtx(2,i2) = derivg(2,i)
                   bmtx(2,i4)=gk2*(-v2(2))
                   bmtx(2, i5) = gk2*v1(2)
                   bmtx(2, i6) = gk2*v3(2)
                   bmtx(3,i3) = derivg(3,i)
                   bmtx(3, i4) = gk3*(-v2(3))
                   bmtx(3, i5) = gk3*v1(3)
                   bmtx(3, i6) = gk3*v3(3)
                   bmtx(4,i1) = derivg(2,i)
                   bmtx(4,i2) = derivg(1,i)
                   bmtx(4,i4)=gk2*(-v2(1))+gk1*(-v2(2))
                   bmtx(4,i5)=gk2*v1(1)+gk1*v1(2)
                   bmtx(4,i6)=gk2*v3(1)+gk1*v3(2)
                   bmtx(5,i2) = derivg(3,i)
                   bmtx(5,i3) = derivg(2,i)
                   bmtx(5,i4)=gk3*(-v2(2))+gk2*(-v2(3))
                   bmtx(5,i5)=gk3*v1(2)+gk2*v1(3)
                   bmtx(5,i6)=gk3*v3(2)+gk2*v3(3)
                   bmtx(6,i1) = derivg(3,i)
                                     96
```

```
bmtx(6,i3) = derivg(1,i)
                  bmtx(6, i4) = gk3*(-v2(1)) + gk1*(-v2(3))
                  bmtx(6, i5) = gk3*v1(1) + gk1*v1(3)
                  bmtx(6, i6) = gk3*v3(1) + gk1*v3(3)
  110
            continue
С
                  do 120 i=1.6
                  do 120 j=1, neldof
  120
                         bmtxt(j,i)=bmtx(i,j)
С
            weight = wx * wy * wz
            detwt = det * weight
c *** calculate strain
            call mmult(6,24,1,bmtx,edisp,estrainp,0,1.0)
c *** Transform strain to local coordinates system
            call mmult(6,6,1,rot6,estrainp,estrain,0,1.0)
c *** Subtract any prior plastic strain
            do 130 i=1,6
                  estrain(i) = estrain(i) - stssh4(12+i,issts)
  130
c *** Calculate stress using plane-strain formulas
      estress(1) = elast/(1.0-poiss**2)*(estrain(1) + poiss*estrain(2))
      estress(2) = elast/(1.0-poiss**2)*(poiss*estrain(1) + estrain(2))
                  estress(3) = elast * estrain(3)
                  estress(4) = shear * estrain(4)
                 estress(5) = 5.0 / 6.0 * shear * estrain(5)
                  estress(6) = 5.0 / 6.0 * shear * estrain(6)
c *** Transform stress back to global coordinates
С
            call mmult(6,6,1,rot6t,estress,estressg,0,1.0)
c *** find eforceh for this gauss point and sum to element force
C
            call mmult(24,6,1,bmtxt,estressg,eforceh,1,detwt)
  140
            continue
            endif
 *** end of hourglass loop
С
c *** numerical integration loop for membrane
            icount=0
            do 300 ix=1,ngausx
             rc=gausx(ix)
             wx=weighx(ix)
             do 300 iy=1,ngausy
              sc=gausy(iy)
              wy=weighy(iy)
              do 300 iz=1, ngausz
               tc=gausz(iz)
```

```
wz=weighz(iz)
      *** increase counter
C
                                            igausp=igausp+1
                                            issts=(ie-1)*8+igausp
                  Initialize stssh4 if first time
C
                                            if (itime .eq. 1) then
                                                                 do 150 i=13.21
                                                                                           stssh4(i,issts) = 0.0
       150
С
                                                                 stssh4(19,issts) = Syp
                                                                 stssh4(20,issts) = Phi0
                                           endif
      *** compute shape functions and derivatives
                                                                 call shp2d4 (rc,sc,shapef)
                                                                 call der2d4 (rc,sc,derivl)
                                                                 call shp1d2 (tc,shape1)
                                                                 call der1d2 (tc,deriv1)
      *** compute jacobian matrix and its inverse
                                                                 hz=thick*0.5*(shape1(2)*(1.0-ref)-shape1(1)*(1.0+ref))
                                                                 hzdt=thick*0.5
                     aj(1,1) = derivl(1,1) *xc(1) + derivl(1,2) *xc(2) + derivl(1,3) *xc(3) +
                                                                                       derivl(1,4)*xc(4)+derivl(1,1)*hz*v3(1)+
                 1
                                                                                       derivl(1,2)*hz*v3(1)+derivl(1,3)*hz*v3(1)+
                 2
                                                                                      derivl(1,4)*hz*v3(1)
                 3
                     aj(2,1) = derivl(2,1) *xc(1) + derivl(2,2) *xc(2) + derivl(2,3) *xc(3) +
                                                                                       derivl(2,4)*xc(4)+derivl(2,1)*hz*v3(1)+
                 1
                                                                                       derivl(2,2)*hz*v3(1)+derivl(2,3)*hz*v3(1)+
                 2
                 3
                                                                                       derivl(2,4)*hz*v3(1)
                                                                 ai(3,1) = hzdt *v3(1)
                    aj(1,2) = derivl(1,1) *yc(1) + derivl(1,2) *yc(2) + derivl(1,3) *yc(3) + derivl(1,3) *yc(3)
                                                                                       derivl(1,4)*yc(4)+derivl(1,1)*hz*v3(2)+
                 1
                                                                                       derivl(1,2)*hz*v3(2)+derivl(1,3)*hz*v3(2)+
                 2
                 3
                                                                                       derivl(1,4)*hz*v3(2)
                    aj(2,2) = derivl(2,1) *yc(1) + derivl(2,2) *yc(2) + derivl(2,3) *yc(3) + derivl(3,3) *yc(3,3) *yc(3) + derivl(3,3) *yc(3) + derivl(3,3) *yc(3) + derivl(3,
                                                                                       derivl(2,4)*yc(4)+derivl(2,1)*hz*v3(2)+
                 1
                                                                                       deriv1(2,2)*hz*v3(2)+deriv1(2,3)*hz*v3(2)+
                 2
                                                                                       derivl(2,4)*hz*v3(2)
                 3
                                                                 ai(3,2) = hzdt * v3(2)
                     aj(1,3) = derivl(1,1)*zc(1)+derivl(1,2)*zc(2)+derivl(1,3)*zc(3)+
                                                                                       derivl(1,4)*zc(4)+derivl(1,1)*hz*v3(3)+
                                                                                       derivl(1,2)*hz*v3(3)+derivl(1,3)*hz*v3(3)+
                 2
                                                                                       derivl(1,4)*hz*v3(3)
                 3
                     aj(2,3) = derivl(2,1)*zc(1)+derivl(2,2)*zc(2)+derivl(2,3)*zc(3)+.
                                                                                       derivl(2,4)*zc(4)+derivl(2,1)*hz*v3(3)+
                 1
                                                                                       derivl(2,2)*hz*v3(3)+derivl(2,3)*hz*v3(3)+
                 2
                                                                                       derivl(2,4)*hz*v3(3)
                 3
```

```
ai(3,3) = hzdt * v3(3)
C
                    call inv3x3 (aj,ajinv,det)
С
      compute global derivatives and strain-nodal displacement matrix
С
                   do 160 i=1, n
      derivg(1,i)=ajinv(1,1)*derivl(1,i)+ajinv(1,2)*derivl(2,i)
      derivg(2,i) = ajinv(2,1)*derivl(1,i)+ajinv(2,2)*derivl(2,i)
      derivg(3,i) = ajinv(3,1) * derivl(1,i) + ajinv(3,2) * derivl(2,i)
                   derilg(1,i)=ajinv(1,3)*hzdt
                    derilg(2,i)=ajinv(2,3)*hzdt
                    derilg(3,i)=ajinv(3,3)*hzdt
             continue
  160
С
                    do 170 i=1,6
                   do 170 j=1,24
                          bmtx(i,j)=0.0
             continue
  170
                   do 180 i=1,n
                    i1=(i-1)*6+1
                    i2=i1+1
                    i3=i2+1
                    i4=i3+1
                    i5=i4+1
                    i6=i5+1
                    gkl=derivg(1,i)*hz+shapef(i)*derilg(1,i)
                    gk2=derivg(2,i)*hz+shapef(i)*derilg(2,i)
                    gk3=derivg(3,i)*hz+shapef(i)*derilg(3,i)
С
                   bmtx(1,i1) = derivg(1,i)
                   bmtx(1,i4)=gk1*(-v2(1))
                   bmtx(1, i5) = gk1*v1(1)
                   bmtx(1,i6) = gk1*v3(1)
                   bmtx(2,i2) = derivg(2,i)
                   bmtx(2,i4)=gk2*(-v2(2))
                   bmtx(2, i5) = gk2*v1(2)
                   bmtx(2, i6) = gk2*v3(2)
                   bmtx(3,i3) = derivg(3,i)
                   bmtx(3,i4)=gk3*(-v2(3))
                   bmtx(3, i5) = gk3*v1(3)
                    bmtx(3, i6) = gk3*v3(3)
                   bmtx(4,i1) = derivg(2,i)
                    bmtx(4,i2) = derivg(1,i)
                    bmtx(4,i4)=gk2*(-v2(1))+gk1*(-v2(2))
                    bmtx(4,i5) = gk2*v1(1) + gk1*v1(2)
                    bmtx(4,i6) = gk2*v3(1) + gk1*v3(2)
                    bmtx(5,i2) = derivg(3,i)
                    bmtx(5,i3) = derivg(2,i)
                    bmtx(5,i4)=gk3*(-v2(2))+gk2*(-v2(3))
                    bmtx(5, i5) = gk3*v1(2) + gk2*v1(3)
                    bmtx(5, i6) = gk3*v3(2) + gk2*v3(3)
                    bmtx(6,i1) = derivg(3,i)
                    bmtx(6,i3) = derivg(1,i)
                    bmtx(6,i4)=gk3*(-v2(1))+gk1*(-v2(3))
                    bmtx(6, i5) = gk3*v1(1) + gk1*v1(3)
                    bmtx(6, i6) = gk3*v3(1) + gk1*v3(3)
  180
             continue
```

```
c *** calculate transpose of bmtx
                  do 190 i=1,6
                  do 190 j=1, neldof
                         bmtxt(j,i)=bmtx(i,j)
  190
С
            weight = wx * wy * wz
                  detwt = det * weight
                  icount=icount+1
                  tmp(icount)=detwt
C
c *** calculate strain
С
            call mmult(6,24,1,bmtx,edisp,estrainp,0,1.0)
C
c *** Transform strain to local coordinates system
            call mmult(6,6,1,rot6,estrainp,estrain,0,1.0)
С
c *** Subtract any prior plastic strain
            do 200 i=1.6
                  estrain(i) = estrain(i) - stssh4(12+i,issts)
  200
С
            stress0 = stssh4(19, issts)
            Phi = stssh4(20, issts)
            plastrn = stssh4(21,issts)
c *** Calculate stress using plane-strain formulas
      estress(1) = elast/(1.0-poiss**2)*(estrain(1) + poiss*estrain(2))
      estress(2) = elast/(1.0-poiss**2)*(poiss*estrain(1) + estrain(2))
                  estress(3) = elast * estrain(3)
                  estress(4) = shear * estrain(4)
                  estress(5) = 5.0 / 6.0 * shear * estrain(5)
                  estress(6) = 5.0 / 6.0 * shear * estrain(6)
c *** Elastic-Plastic Calculations
            p = -(estress(1) + estress(2) + estress(3)) / 3.0
C
            do 210 i=1,6
                        s(i) = estress(i) + p * delt(i)
  210
            q = sqrt(1.5 *(s(1)**2 + s(2)**2 + s(3)**2 + 2.0 * s(4)**2 +
                  2.0 * s(5)**2 + 2.0 * s(6)**2))
            F = (q / stress0)**2+2.0*q1*Phi*cosh(-1.5*q2*p/stress0)-
                  (1.0+q3* Phi**2)
С
             if (F.gt.0) then
                  iter = 0
                  dep = 0.0
                  deq = 0.0
c ** Begin Iteration Loop **
                  c1 = 3.0*q2/(2.0*stress0**2)
  220
                  Fq1 = 2.0 * q / (stress0**2)
                  Fq2 = 2.0 / (stress0**2)
                                   100
```

```
Fp1 = 2.0 * q1 * Phi * -c1 * sinh(-c1 * p)
Fp2 = 2.0 * q1 * Phi * c1**2 * cosh(-c1 * p)
                 Fs1 = -q **2 / (2.0 * stress0**3) + 2.0*q1*Phi*c1*
                        p*sinh(-c1*p)
                 Fsp = 2.0*q1*Phi*(-c1**2 * p*cosh(-c1*p) +
    1
                 c1/stress0 * sinh(-c1*p))
                 Fsq = -q / stress0**3
                 stress1 = Syp
                 es1 = stress0/elast
                 e1 = -p*dep/((1-Phi)*stress0)
                 ps1 = plastrn + e1 + es1
                 iflag = 0
                 do 230 i=1,iseg
                        if (iflag .eq. 0) then
                         if(psl .lt. strain(i+1)) then
                          stress1 = stress1 + slope(i)*(ps1-strain(i))
                          iseg1 = i
                          iflag = 1
                         else
                    stress1 = stress1 + slope(i)*(strain(i+1)-strain(i))
                         endif
                        endif
230
                 continue
```

```
С
        Numerical Error Trap
С
                   if ((stress1 .eq. stress0) .or. (abs(e1) .eq. 0.0))
then
                         dsdep = slope(iseg1)
                   else
                         dsdep = (stress1-stress0)/e1
                   endif
                   if (abs(dsdep) .gt. elast) then
                         dsdep = slope(iseg1)
                   endif
С
                   stress1 = Syp
                  el = q * deq / ((1-Phi) * stress0)
                  ps1 = plastrn + e1
                   iflag = 0
                  do 240 i=1, iseq
                    if (iflag .eq. 0) then
                     if(ps1 .lt. strain(i+1)) then
                      stress1 = stress1 + slope(i)*(ps1-strain(i))
                      iseg1 = i
                      iflag = 1
                     else
                     stress1 = stress1 + slope(i)*(strain(i+1)-
strain(i))
                     endif
                    endif
  240
                  continue
С
c *** Numerical Error Trap
                  if ((stress1 .eq. stress0).or.(abs(e1) .eq. 0.0)) then
                         dsdeq = slope(iseg1)
                  else
                         dsdeq = (stress1-stress0)/e1
                  endif
                  if (abs(dsdeq) .gt. elast) then
                         dsdeq = slope(iseg1)
                  endif
c *** Using only stress0 derivatives
                  A(1,1) = bulk * Fp1+Fs1*dsdep
                  A(1,2) = -3.0*shear*Fq1+Fs1*dsdeq
                  A(2,1) = Fq1+deq*(bulk*Fp2+Fsp*dsdep)+dep*Fsq*dsdep
            A(2,2) = Fp1+dep*(-3.0*shear*Fq2+Fsq*dsdeq)+deq*Fsp*dsdeq
С
                  B(1) = -1.0 * F
                  B(2) = -1.0 * dep * Fq1 - deq * Fp1
C
                  call inv2x2(A,Ainv,d)
С
                  call mmult (2, 2, 1, Ainv, B, corr, 0, 1.0)
C
                  Phi = stssh4(20, issts)
                  plastrn = stssh4(21,issts)
                  stress0 = stssh4(19,issts)
C
                  dep = dep + corr(1)
                                    102
```

```
deq = deq + corr(2)
С
                  degrowth = 0.0
                  do 250 i=1,5
                         ind = iplane(i)
                         dstrnplst(ind) = dep * delt(ind) / 3.0 +
                         deg * 3.0 * s(ind) / (2.0 * q)
  250
                  continue
                  dstrnplst(3) = dep / 3.0 + deq * estress(3) / q
С
                  do 260 i=1,3
  260
                        degrowth = degrowth + dstrnplst(i)
С
c ** Calculate New stress, p, q, and F
      estress(1) = elast/(1.0-poiss**2)*(estrain(1) - dstrnplst(1) +
    1
                   poiss*(estrain(2)-dstrnplst(2)))
                   estress(2) = elast/(1.0-poiss**2)*
      1
                (poiss*(estrain(1)-dstrnplst(1))+estrain(2)-dstrnplst(2))
                  estress(3) = elast * (estrain(3)-dstrnplst(3))
                  estress(4) = shear * (estrain(4)-dstrnplst(4))
            estress(5) = 5.0 / 6.0 * shear * (estrain(5)-dstrnplst(5))
            estress(6) = 5.0 / 6.0 * shear * (estrain(6)-dstrnplst(6))
                  p = (estress(1) + estress(2) + estress(3)) / -3.0
С
                  do 270 i=1,6
                         s(i) = estress(i) + p * delt(i)
  270
C
                  deltep = (q*deq - p * dep)/((1-Phi)*stress0)
c Prevent reduction in plastic strain
                  if (deltep .lt. 0.0) then
                        deltep = 0.0
                  endif
С
                  plastrn = plastrn + deltep
                  if (fn .ne. 0.0) then
             anucl = fn/(sn*2.50662827463)*exp(-.5*((plastrn-en)/sn)**2)
                   anucl = 0.0
                  endif
С
                  Phil = Phi + (1 - Phi) * degrowth + anucl * deltep
c Phi cannot decrease
                  if (Phil .gt. Phi) Phi = Phil
c Turn off Void Effects if q1 = 0.0
                  if (q1 .eq. 0.0) Phi = 0.0
c Calculate Strain Hardening
                  elastrn = stress0/elast
                  stress0 = Syp
                  iflag = 0
                  do 280 i=1, iseg
                        if (iflag .eq. 0) then
                         ps1 = plastrn + elastrn
                         if(ps1 .lt. strain(i+1)) then
                           stress0 = stress0 + slope(i)*(psl-strain(i))
                                   103
```

```
iflag = 1
                          else
                    stress0 = stress0 + slope(i)*(strain(i+1)-strain(i))
                         endif
                        endif
  280
                  continue
С
            q = sqrt(1.5 *(s(1)**2 + s(2)**2 + s(3)**2 + 2.0 * s(4)**2 +
                        2.0 * s(5)**2 + 2.0 * s(6)**2)
C
            F = (q / stress0)**2+2.0*q1*Phi*cosh(-1.5*q2*p/stress0)-
                         (1.0+q3* Phi**2)
      1
C
                  iter = iter + 1
C
                  if (iter.gt.500) then
                        write(*,*) 'Failed to Converge'
                        stop
                  endif
                  if(abs(F).gt.eps) goto 220
C
            endif
c *** End of Plastic Deformation Section
C
C
c *** Transform stress back to global coordinates
            call mmult(6,6,1,rot6t,estress,estressg,0,1.0)
c *** find eforce for this gauss point and sum to element force
            call mmult(24,6,1,bmtxt,estressg,eforce,1,detwt)
C
c *** save stress and strain
            do 290 i = 1,6
                  stssh4(i,issts)=estress(i)
                  stssh4(i+6,issts)=estrain(i) - dstrnplst(i)
                  stssh4(i+12,issts)=stssh4(i+12,issts) + dstrnplst(i)
  290
            continue
C
            stssh4(19,issts) = stress0
            stssh4(20,issts) = Phi
            stssh4(21,issts) = plastrn
c *** end of integration loop
  300
            continue
c *** Hourglass Control Correction
С
      do 310 i = 1,24
            eforce(i) = eforce(i) + hour * (eforceh(i) - eforce(i))
  310
c *** Transform Moments back to Global Coordinate system
            do 320 i=1.4
            do 320 j=1,3
                        eforceg((i-1)*6+j)=eforce((i-1)*6+j)
                  eforceg((i-1)*6+3+j)=eforce((i-1)*6+4)*t1(j,1)
                                   104
```

```
+eforce((i-1)*6+5)*t1(j,2)
                                     +eforce((i-1)*6+6)*t1(j,3)
  320
             continue
С
c *** compute lumped mass matrix
С
             sum=0.0
             do 340 i=1, ngauss
  340
             sum=sum+tmp(i)
С
       ems=sum/n
С
             do 350 i=1, neldof
  350
            emass(i)=ems*densty
С
c *** assemble mass and force matrices into system matrix
             do 360 i=1, neldof
             sforce(lindex(i)) = sforce(lindex(i)) + eforceg(i)
             smass(lindex(i)) = smass(lindex(i)) + emass(i)
  360
             continue
С
c *** end of element loop
  370 continue
         return
         end
```

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